



(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date
23 December 2004 (23.12.2004)

PCT

(10) International Publication Number
WO 2004/111183 A2

(51) International Patent Classification⁷:

C12N

(21) International Application Number:

PCT/IL2004/000549

(22) International Filing Date: 20 June 2004 (20.06.2004)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

60/479,467 19 June 2003 (19.06.2003) US

(71) Applicant (*for all designated States except US*): EVO-GENE LTD. [IL/IL]; 13 Gad Finstein Street, 76 121 Rehovot (IL).

(72) Inventors; and

(75) Inventors/Applicants (*for US only*): RONEN, Gil [IL/IL]; 20 Moshav Ometz, 33 870 Emeq Hefer (IL). RABINOVICH, Larisa [IL/IL]; 13/13 Hanagid Street, 75 482 Rishon LeZion (IL). MEISSNER, Rafael [IL/IL]; 11 Halevona Street, 76 350 Rehovot (IL). KARCHI, Hagai [IL/IL]; Moshav Sitiya, 76 834 Moshav Sitiya (IL).

(74) Agent: G.E. EHRLICH (1995) LTD.; 11 Menachem Begin Street, 52 521 Ramat Gan (IL).

(81) Designated States (*unless otherwise indicated, for every kind of national protection available*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(84) Designated States (*unless otherwise indicated, for every kind of regional protection available*): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

- without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

WO 2004/111183 A2

(54) Title: NUCLEOTIDE SEQUENCES FOR REGULATING GENE EXPRESSION IN PLANT TRICHOMES AND CONSTRUCTS AND METHODS UTILIZING SAME

(57) Abstract: Novel plant derived regulatory sequences and constructs and methods of using such sequences for directing expression of exogenous polynucleotide sequences in trichomes are provided.

NUCLEOTIDE SEQUENCES FOR REGULATING GENE EXPRESSION IN
PLANT TRICHOMES AND CONSTRUCTS AND METHODS UTILIZING SAME

5 FIELD AND BACKGROUND OF THE INVENTION

The present invention relates to nucleotide sequences for regulating gene expression in plant trichomes and, more particularly, to methods of utilizing such nucleotide sequences for synthesizing polypeptides and molecules of interest in plant trichomes.

10 Polypeptides can be expressed in a wide variety of cellular hosts. For economic reasons, genetically engineered unicellular microorganisms are most widely used for commercial production of polypeptides. However, in some cases, expression of mammalian proteins in unicellular organisms results in incorrect folding and processing of the expressed polypeptides leading to loss of biological or
15 physiological activity of the obtained polypeptide. For these reasons, attempts have been made, with varying degrees of success, to express mammalian polypeptides in plants.

Transgenic plants are fast becoming a preferred system for the expression of many recombinant proteins, especially those intended for therapeutic purposes. One
20 advantage of using plants is the potential for protein production on an agricultural scale at an extremely competitive cost. Among other advantages is that most plant transformation techniques result in a stable integration of the foreign DNA into the plant genome, so genetic recombination by crossing of transgenic plants is a simple method for introducing new genes, accumulating multiple genes into plants and
25 avoiding the contamination of pathogens such as viruses and prions, which may affect human and animals. Furthermore, the processing and assembly of recombinant proteins in plants may also complement that in mammalian cells, which may be an advantage over the more commonly used microbial expression systems.

Although plants provide a suitable alternative to unicellular expression systems, several disadvantages characterize current approaches for production of protein in plants. First, the concentration of the produced protein is typically low (around 1% of total proteins) making purification extremely difficult. Second, other compounds may interfere with protein purification or even damage the proteins during purification. Third, expressing foreign proteins in propagated plants can lead

to environmental contamination and health risks associated with unwanted production of those proteins in cross pollinated plants.

In efforts of overcoming the above described limitations and while reducing the present invention to practice, the present inventors have discovered that plant trichomes enable compartmentalized production of foreign proteins as well as enzymatic production of novel chemicals, since many types of chemicals are naturally produced and even secreted from trichomes.

The above ground surfaces of many plants are covered with trichomes or hairs. The morphology of these structures can vary greatly with tissue type and species. Indeed, the botanical literature contains more than 300 descriptions (uniseriate, capitate-sessile, etc.) of various morphological types of such hairs (3, and references therein). Functionally, trichomes may be simple hairs that deter herbivores, guide the path of pollinators, or affect photosynthesis, leaf temperature, or water loss through increased light reflectance as in desert species. Alternatively, they may be more specialized tissues (glandular secreting trichomes) whose principal function(s) may be to produce pest- or pollinator-interactive compounds that are stored or volatilized at the plant surface. It has been suggested that in some desert species the principal role of glandular secreting trichomes is to produce such high levels of exudate that it forms a continuous layer on the plant surface. This layer may increase light reflectance and thereby reduce leaf temperature (30).

Trichomes develop projections from protodermal cells. Their structures arise from a series of anticlinal and periclinal divisions to form supporting auxiliary cells and glands. The appearance of glands atop supporting cells and the occurrence of exudate around gland cells has suggested that secretions are produced in gland cells and not by other epidermal or subepidermal cells.

In several species, such as tomato and potato, a unique type of trichomes accumulates certain protein (polyphenol oxidase) and compound (polyphenol) in the associated glands on the top of the trichome. When an insect lands on a leaf surface and contacts these trichomes, they discharge their inner compounds thereby contacting the insect and smearing it with a brown sticky compound, which is the product of enzymatic oxidation of the polyphenols (reviewed in 4).

The mass production, accumulation, and secretion of such proteins and chemicals involve a specific genetic mechanism. This genetic mechanism includes

genes (5, 6) and promoters (7, 8, 9) acting in trichome cells and cells organelles suited for accumulation and secretion of mass products. This genetic mechanism allows, for example, trichome exudates to reach 16% of total dry weight of leaves of a certain tobacco species (10) and a single protein to reach 60% of total proteins or a concentration of 14 mg/mL in the trichome content of a solanum species (11, 12). The use of this genetic mechanism was suggested for tissue specific production and accumulation of natural and heterologous proteins as well as chemicals (6). New compounds produced can be beneficial for the plant itself by increasing resistance against pests such as insects, bacteria, and fungi (6), or for Molecular Farming or Bio-Farming of human or mammalian proteins for the use as therapeutics. In the latter, harvesting the proteins produced in the trichomes can be mechanized.

Directing protein expression into trichome cells may involve the use of polynucleotides originated form different origins. A candidate source for such regulatory elements is cotton as its fiber tissue is structurally modified trichomes. The promoter sequences of cotton fiber specific genes were shown to direct β -glucuronidase (GUS) expression to the trichome cells of tobacco plants (7, 9). Alteration of trichomes structure or chemistry by, for example, increasing cotton trichome length or by producing pigments in the fiber could be beneficial for the cotton industry.

Natural chemicals of trichomes are already used as flavor, aroma, medicinals, pesticides, and cosmetic ingredients (13, 14). Natural chemicals content was altered using antisense and co-suppression methods (6). However, enzymatic modifications of trichomes compounds via genetic engineering of genes, designed to produce other useful compounds in trichomes, was never shown before.

Several limitations had narrowed so far the use of plant trichomes for commercially production of heterologous proteins and novel chemicals.

First, protein yield is very limited in trichome cells and to date there is no existing method that enables commercially significant production of proteins in these cells. Although there are known promoter sequences that are capable of directing protein synthesis in trichomes (7, 8, 9), proteins expressed therefrom accumulated at average levels of accumulation of a single trichome protein and thus these promoters cannot be considered commercially useful for protein production, as is. Second, trichomes usually produce a mix of several metabolites, some of which (e.g., phenols

and alkaloids), can inhibit protein accumulation or substantially hinder purification of desired compounds produced in trichomes (See material and methods in 12). Thus, reducing the levels of such harmful metabolites is required in order to improve harvesting and collection of the desired products. Third, the production of novel 5 compounds in plants always involves risks of escape of genetic material (pollen and seeds) to the environment with potential damage to other organisms (plants, insects animals, human). Hence, when producing novel compounds one should consider the elimination of the possible spread of the new genetic material.

There is thus a widely recognized need for and it would be highly 10 advantageous to have nucleotide sequences for regulating gene expression in plant trichomes methods of utilizing such nucleotide sequences for generating molecules of interest in plant trichomes.

SUMMARY OF THE INVENTION

15 According to one aspect of the present invention there is provided an isolated polynucleotide comprising a nucleic acid sequence at least 80 % identical to SEQ ID NO: 23, 26 or 29, wherein the nucleic acid sequence is capable of regulating expression of at least one polynucleotide sequence operably linked thereto in trichomes.

20 According to another aspect of the present invention there is provided a nucleic acid construct comprising the isolated polynucleotide.

According to further features in preferred embodiments of the invention described below, the nucleic acid construct further comprising at least one heterologous polynucleotide operably linked to the isolated polynucleotide.

25 According to still further features in the described preferred embodiments the nucleic acid construct further comprises, a nucleic acid sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a subcellular compartment of a trichome.

According to still further features in the described preferred embodiments the 30 nucleic acid sequence is selected from the group consisting of SEQ ID NOS: 59, 61, 63, 65 and 67.

According to yet another aspect of the present invention there is provided a transgenic cell comprising the nucleic acid construct.

According to still another aspect of the present invention there is provided a transgenic plant comprising the nucleic acid construct.

According to an additional aspect of the present invention there is provided an isolated polynucleotide comprising a nucleic acid sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a subcellular compartment of a trichome, wherein the peptide is encoded by the polynucleotide sequence set forth in SEQ ID NO: SEQ ID NOS: 59, 61, 63, 65 and 67.

According to yet an additional aspect of the present invention there is provided a nucleic acid construct comprising the isolated polynucleotide.

According to still further features in the described preferred embodiments the nucleic acid construct further comprising an expressible polynucleotide sequence translationally fused to the nucleic acid sequence encoding the peptide.

According to still an additional aspect of the present invention there is provided a method of producing a polypeptide of interest in plant trichomes, the method comprising:

- (a) expressing the polypeptide of interest in the plant trichomes; and
- (b) down-regulating a level of at least one molecule endogenous to the plant trichomes, the at least one molecule being capable of interfering with expression, accumulation or stability of the polypeptide of interest.

According to still further features in the described preferred embodiments step (b) is effected by gene silencing.

According to a further aspect of the present invention there is provided a method of producing a molecule of interest in plant trichomes, the method comprising:

(a) expressing a polypeptide capable of directly or indirectly increasing a level of the molecule of interest in the plant trichomes; and (b) down-regulating a level of at least one molecule endogenous to the plant trichomes, the at least one molecule being capable of interfering with accumulation or stability of the molecule of interest, thereby producing the molecule in the plant trichomes.

According to still further features in the described preferred embodiments the polypeptide is endogenously expressed in the plant trichomes.

According to still further features in the described preferred embodiments the expressing the polypeptide in the plant trichomes is effected by introducing into the

plant trichomes a nucleic acid sequence encoding the polypeptide positioned under a transcriptional control of a promoter functional in the plant trichomes.

According to still further features in the described preferred embodiments the promoter is as set forth in SEQ ID NO: 23, 26, 29, 35, 38, 39, 42, 48, 50 or 51.

5 According to still further features in the described preferred embodiments the nucleic acid sequence encoding the polypeptide of interest further encodes a peptide capable of directing transport of the polypeptide fused thereto into a subcellular compartment of the plant trichomes.

According to still further features in the described preferred embodiments the
10 at least one molecule is an enzyme or a metabolite.

According to still further features in the described preferred embodiments the metabolite is selected from the group consisting of polyphenols, ketones, terpenoids, phenylpropanoids and alkaloids.

According to still further features in the described preferred embodiments the
15 enzyme is PPO.

According to still further features in the described preferred embodiments step (b) is effected by gene silencing.

According to yet a further aspect of the present invention there is provided a plant genetically modified to express a molecule of interest in trichomes, wherein the
20 plant is further modified or selected capable of accumulating less than 50 % of average volume of undesired compounds in trichome cells of the plant species.

According to still further features in the described preferred embodiments at least a portion of cells of the plant are genetically modified to include an expression construct including a polynucleotide sequence of a trichome specific promoter.

25 According to still further features in the described preferred embodiments the expression construct further includes an additional polynucleotide sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a subcellular compartment of the trichome, whereas the additional polynucleotide is translationally fused to the polynucleotide sequence.

30 According to still further features in the described preferred embodiments at least a portion of cells of the plant are genetically modified to include an expression construct including a first polynucleotide sequence encoding the polypeptide

translationally fused to a second polynucleotide sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a trichome.

According to still further features in the described preferred embodiments the expression or accumulation is in a subcellular compartment of trichomes.

5 According to still further features in the described preferred embodiments the subcellular compartment is a leucoplast.

According to still further features in the described preferred embodiments the trichome specific promoter is set forth by SEQ ID NO: 23, 26 or 29.

10 According to still further features in the described preferred embodiments the trichome specific promoter is set forth by SEQ ID NO: 23, 26, 29, 35, 38, 39, 42 or 45.

According to still further features in the described preferred embodiments the additional polynucleotide sequence is set forth by SEQ ID NO: 59, 61, 63, 65 or 67.

15 40. The plant of claim 31, wherein the molecule of interest is not a reporter polypeptide.

According to still further features in the described preferred embodiments the plant is modified or selected capable of generating a trichome density above 50,000 trichomes/gr leaf tissue.

20 According to still further features in the described preferred embodiments the plant is modified or selected capable of generating a trichome size of 50 % above average size of the plant species.

According to still further features in the described preferred embodiments the plant is modified or selected capable of generating leaf surface size at least 25 % above average size of the plant species.

25 According to still further features in the described preferred embodiments the plant is modified or selected capable of generating total leaf number at least 50 % above average leaf number of the plant species.

According to still further features in the described preferred embodiments the plant is sterile.

30 According to still further features in the described preferred embodiments the plant is further genetically modified capable of secreting the exogenous polypeptide from trichome cells.

According to still a further aspect of the present invention there is provided a method of harvesting trichomes and/or exudates and/or content thereof, the method comprising: (a) incubating a trichome-containing plant tissue in a liquid such that trichome exudates and content is released into the liquid, wherein incubating is effected while avoiding friction of the trichome-containing plant tissue with a solid phase; and (b) collecting the liquid, to thereby harvest the trichome exudates and content.

According to still further features in the described preferred embodiments the liquid includes an antioxidant.

According to still further features in the described preferred embodiments the antioxidant is selected from the group consisting of citric acid, ascorbic acid and sodium bisulfite

According to still further features in the described preferred embodiments the liquid is water.

According to still further features in the described preferred embodiments the trichome-containing plant tissue is selected from the group consisting of a shoot, a flower and a leaf.

According to still a further aspect of the present invention there is provided an apparatus for mechanical harvesting of trichome exudates and content, the apparatus comprising a mechanism designed and configured for mechanically agitating a trichome-containing plant tissue in a fluid and collecting the fluid to containing the trichome exudates or content..

The present invention successfully addresses the shortcomings of the presently known configurations by providing nucleotide sequence for regulating gene expression in plant trichomes and methods of utilizing such nucleotide sequences for generating molecules in plant trichomes

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, suitable methods and materials are described below. In case of conflict, the patent specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

BRIEF DESCRIPTION OF THE DRAWINGS

The invention is herein described, by way of example only, with reference to the accompanying drawings. With specific reference now to the drawings in detail, it is stressed that the particulars shown are by way of example and for purposes of 5 illustrative discussion of the preferred embodiments of the present invention only, and are presented in the cause of providing what is believed to be the most useful and readily understood description of the principles and conceptual aspects of the invention. In this regard, no attempt is made to show structural details of the invention in more detail than is necessary for a fundamental understanding of the invention, the 10 description taken with the drawings making apparent to those skilled in the art how the several forms of the invention may be embodied in practice.

In the drawings:

FIG. 1a is a prior art schematic illustration of various tomato trichomes. Type VI glandular trichomes naturally accumulate high levels of the PPO enzyme 15 (Luckwill LC. 1943. The Aberdeen University Press, Aberdeen, Scotland).

FIGs. 1b-h are photomicrographs depicting trichome specific expression of GUS under the regulation of the CaMV 35S, TR2, TR5, TR11, TR25 or TR27P promoters. Figure 1b - Trichomes of wild-type tomato plants. Figure 1c - Trichomes of tomato plants overexpressing GUS under the constitutive CaMV 35S promoter. 20 Figures 1b-h - Trichomes of tomato plants overexpressing GUS with different trichome promoters

FIG. 2 is a photograph depicting total protein yield of chemically extracted trichome cells as determined by coomassie staining.

FIG. 3 is a photograph depicting decreased PPO activity in the presence of 25 increasing concentrations of sodium bisulfite, as indicated by medium browning, which is indicative of PPO activity.

FIGs. 4a-c schematically illustrate various embodiments of a trichome mechanical harvester constructed in accordance with the teachings of the present invention.

FIGs. 5a-c are graphs showing the effect of pruning on leaf number of 30 3 tomato cultivars. Figure 5a cultivar 678; Figure 5b cultivar 1312; Figure 5c cultivar 2545. Treatment 1 - Plant shoot number was not limited, plant height was limited to 1 m, flowers were cut-off before fruit set; Treatment 2 - plant shoot number was not

10

limited, plant height was limited to 2 m, flowers were cut-off before fruit set; Treatment 3 - plant shoot number was not limited, leading apical meristem was cut (i.e. breaking apical dominance) when reached 0.5m, flowers were cut-off before fruit set. Treatment 4 – control plants were treated for tomato fruit set, such that each plant 5 includes 2 shoots. Flowers and fruits were untouched.

FIGs. 6a-d are graphs depicting expression levels of three trichome-expressed genes as determined by RT-PCR. Expression is shown as fold increase over house-keeping gene expression. 273_1 is *L.hirsutum* var glabratum cultivar; 247 is *L. esculentum* cultivar. Tissue key: L- Leaves; L-T- Leaves minus Trichomes; T1 and 10 T2 are two independent RNA samples of Trichome cells.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention is nucleotide sequence for regulating gene expression in plant trichomes which can be utilized for generating molecules in plant trichomes. 15 Specifically the present invention is of plants which are modified for enhanced expression and accumulation of molecules in plant trichomes.

The principles and operation of the present invention may be better understood with reference to the drawings and accompanying descriptions.

Before explaining at least one embodiment of the invention in detail, it is to be 20 understood that the invention is not limited in its application to the details set forth in the following description or exemplified by the Examples. The invention is capable of other embodiments or of being practiced or carried out in various ways. Also, it is to be understood that the phraseology and terminology employed herein is for the purpose of description and should not be regarded as limiting.

Trichomes are hairy-like epidermal multi-cell structures found on the outer 25 surface of leaves, stems and flowers of about 20-30 % of plant species. Their Main function is associated with plant protection against insects, microbes and herbivores due to their ability to massively accumulate and secrete pest-deterrant phytochemicals. Other functions include water absorption, seed dispersal and abrasion protection.

The high production capacity of trichomes prompted their utilization as “green-factories” for producing commercially useful compounds (e.g., U.S. Pat. No. 30 6,730,826). However, mass production of recombinant proteins in trichomes is limited

by poor production efficiency and the presence of metabolites and enzymes, which may interfere with purification and activity of the desired compounds.

While reducing the present invention to practice the present inventors have devised a novel approach for increasing expression, accumulation and harvesting of molecules in plant trichomes, while reducing the chances of accidentally spreading the non-plant genetic material used to generate the molecules.

As is illustrated in the Examples section, the present inventors have uncovered that by reducing the concentration of undesired compounds in trichome exudates an enhanced level of expression, accumulation and/or purification of commercially valuable molecules within the trichomes can be achieved. Furthermore the present inventors uncovered through laborious experimentation and time consuming analysis a number of novel trichome active regulatory elements (see Example 1), which enable protein over-expression in trichomes (see Examples 2-4).

These findings allow, for the first time, improved molecular farming in trichome cells.

Thus, according to one aspect of the present invention there is provided a method of producing a molecule of interest in plant trichomes.

As used herein the term "trichome" refers to both a "simple" (also termed "non-glandular") trichome and a "glandular-secreting" (GST) trichome. Preferably, the term trichome refers to a GST trichome.

As used herein the term "molecule" refers to at least one small molecule chemical (e.g., nicotine, flavonoids). Such molecules can be naturally expressed or present in trichomes or can be direct or indirect expression products of heterologous polynucleotides. Examples of molecules which can be produced in trichome cells according to this aspect of the present invention include, but are not limited to, oils, dyes, flavors, biofuels, or industrial biopolymers, pharmaceuticals, nutraceuticals and cosmeceuticals.

As used herein the term "producing" refers to the process of expressing and/or accumulating the molecule in trichome cells. When appropriate, producing may also refer to subsequent steps of purifying the molecule from the trichome cells.

The method, according to this aspect of the present invention, is effected by upregulating expression of a polypeptide capable of directly or indirectly increasing a level of the molecule of interest; and down-regulating a level of at least one molecule

endogenous to the plant trichome, which is capable of interfering with the production of the molecule in the plant trichomes, thereby producing the molecule in the plant trichomes.

Examples of polypeptides capable of directly or indirectly increasing the level 5 of the molecule of interest include include for example, trichome specific transcription factors which promote expression in trichome cells. Alternatively, the polypeptide can be an enzyme participating in a biochemical pathway, which produces the molecule in the trichome.

Expression of polypeptides in plant trichomes according to this aspect of the 10 present invention, may be effected by placing a polynucleotide encoding the polypeptide of interest under the regulation of a cis-acting regulatory element capable of directing expression from the polynucleotide in trichome cells.

As used herein, the phrase "cis acting regulatory element" refers to a polynucleotide sequence, preferably a promoter, which binds a trans acting regulator 15 and regulates the transcription of a coding sequence located downstream thereto in trichome cells.

It will be appreciated that a regulatory sequence is "operably linked" to a coding polynucleotide sequence if it is capable of exerting a regulatory effect on the coding sequence linked thereto. Preferably, the regulatory sequence is positioned 1- 20 500 bp upstream of the ATG codon of the coding nucleic acid sequence, although it will be appreciated that regulatory sequences can also exert their effect when positioned elsewhere with respect to the coding nucleic acid sequence (e.g., within an intron).

A number of trichome active promoters are known in the art which can be used 25 in accordance with the present invention. Examples include, but are not limited to, the CYP71D16 trichome-specific promoter [Wang E. J Exp Bot. (2002) 53(376):1891-7, see U.S. Pat. No. 6,730,826] and the cotton LTP3 and LTP6 promoters (7,9).

Methods of identifying trichome active or specific promoters are well described in Examples 1-3 of the Examples section which follows.

30 As mentioned hereinabove, the present inventors have identified a number of cis-acting regulatory elements, which are capable of regulating transcription of coding nucleic acid sequences operably linked thereto in trichome cells.

Thus the present invention provides an isolated polynucleotide having a nucleic acid sequence at least 80 % identical to SEQ ID NO: 23, 26 or 29, wherein the nucleic acid sequence is capable of regulating expression of at least one polynucleotide sequence operably linked thereto in trichomes.

5 According to other embodiments of this aspect of the present invention the nucleic acid sequence of the present invention is at least 80 % identical to SEQ ID NO: 35, 38, 39, 42, 45, 48, 51 or 54.

10 Preferably, the polynucleotides (promoters) of the present invention are modified to create variations in the molecule sequences such as to enhance their promoting activities, using methods known in the art, such as PCR-based DNA modification, or standard DNA mutagenesis techniques, or by chemically 15 synthesizing the modified polynucleotides.

Accordingly, the sequences set forth in SEQ ID NOs: 23, 26, 29, 35, 38, 39, 42, 45, 48, 51 and 54 may be truncated or deleted and still retain the capacity of 15 directing the transcription of an operably linked DNA sequence in trichomes. The minimal length of a promoter region can be determined by systematically removing sequences from the 5' and 3'-ends of the isolated polynucleotide by standard techniques known in the art, including but not limited to removal of restriction enzyme fragments or digestion with nucleases.

20 In another approach, novel hybrid promoters can be designed or engineered by a number of methods. Many promoters contain upstream sequences which activate, enhance or define the strength and/or specificity of the promoter, such as described, for example, by Atchison [Ann. Rev. Cell Biol. 4:127 (1988)]. T-DNA genes, for example contain "TATA" boxes defining the site of transcription initiation and other 25 upstream elements located upstream of the transcription initiation site modulate transcription levels [Gelvin In: Transgenic Plants (Kung, S.-D. and Us,R., eds, San Diego: Academic Press, pp.49-87, (1988)]. Another chimeric promoter combined a trimer of the octopine synthase (ocs) activator to the mannopine synthase (mas) activator plus promoter and reported an increase in expression of a reporter gene [Min 30 Ni *et al.*, The Plant Journal 7:661 (1995)]. The upstream regulatory sequences of the present invention can be used for the construction of such chimeric or hybrid promoters. Methods for construction of variant promoters include, but are not limited to, combining control elements of different promoters or duplicating portions or

regions of a promoter (see for example, U.S. Pat. Nos. 5,110,732 and 5,097,025). Those of skill in the art are familiar with the specific conditions and procedures for the construction, manipulation and isolation of macromolecules (e.g., DNA molecules, plasmids, etc.), generation of recombinant organisms and the screening and isolation of genes, [see for example Sambrook *et al.*, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press, (1989); Mailga *et al.*, Methods in Plant Molecular Biology, Cold Spring Harbor Press, (1995); Birren *et al.*, Genome Analysis: volume 1, Analyzing DNA, (1997); volume 2, Detecting Genes, (1998); volume 3, Cloning Systems, (1999); and volume 4, Mapping Genomes, (1999), Cold 5 Spring Harbor, N.Y].

The above-described nucleic acid sequences (promoters) can be used to drive expression of a heterologous polynucleotide of interest in trichome cells. Preferably, the heterologous polynucleotide can encode any naturally occurring or man-made recombinant protein, such as pharmaceutical proteins [e.g., growth factors and 15 antibodies Schillberg Naturwissenschaften. (2003) Apr;90(4):145-55] and food additives. It will be appreciated that molecular farming is a well-proven way of producing a range of recombinant proteins, as described in details in Ma Nat Rev Genet. 2003 Oct;4(10):794-805; Twyman Trends Biotechnol. 2003 Dec;21(12):570-8.

To facilitate accumulation of the polypeptide of interest in trichome cells, it 20 may be beneficial to translationally link the heterologous polynucleotide encoding the polypeptide to a signal peptide-encoding sequence which is capable of directing transport of the polypeptide into sub-cellular organelle of a the trichome. Examples of subcellular organelles of trichome cells include, but are not limited to, leucoplasts, chloroplasts, chromoplasts, mitochondria, nuclei, peroxisomes, endoplasmic reticulum 25 and vacuoles. Preferably the signal peptide of this aspect of the present invention is a leucoplast localization signal. It is appreciated that since the protein is not accumulated in the cytoplasm, but rather in the subcellular organelle of the trichomes, it is expected to be stored in relatively high concentrations without being exposed to the degrading compounds present in the trichome vacuole. Examples of signal 30 peptides which may be used in accordance with the present invention include, but are not limited to, the stroma or lumen directing signal peptides of PPOA and PPOD (SEQ ID NO: 60, 62, 64, 66 and 76, see Example 3). Polynucleotides encoding these signal peptides are set forth in SEQ ID NOs: 59, 61, 63, 65 and 75.

The polynucleotides (i.e., trichome active promoter sequence, signal peptide encoding polynucleotide) of the present invention, or fragments, variants or derivatives thereof, can be incorporated into nucleic acid constructs, preferably expression constructs (i.e., expression vectors), which can be introduced and replicate in a plant cell, such as a trichome. Such nucleic acid constructs may include the heterologous polynucleotide of interest such as described hereinabove, operably linked to any of the promoter sequences of the present invention.

The nucleic acid construct can be, for example, a plasmid, a bacmid, a phagemid, a cosmid, a phage, a virus or an artificial chromosome. Preferably, the nucleic acid construct of the present invention is a plasmid vector, more preferably a binary vector.

The phrase "binary vector" refers to an expression vector which carries a modified T-region from Ti plasmid, allowing multiplication both in *E. coli* and in *Agrobacterium* cells, and usually comprising selection gene(s). Such a binary vector suitable for the present invention is described in Example 1 of the Examples section which follows.

The nucleic acid construct of the present invention can be utilized to transform a host cell. Preferably a plant cell. Preferably, the nucleic acid construct of the present invention is used to transform at least a portion of cells of a plant.'

Methods of introducing nucleic acid constructs into a cell or a plant are well known in the art. Accordingly, suitable methods for introducing nucleic acid sequences into plants include, but are not limited to, bacterial infection, direct delivery of DNA (e.g., via PEG-mediated transformation, desiccation/inhibition-mediated DNA uptake, electroporation, agitation with silicon carbide fibers, and acceleration of DNA coated particles, such as described by Potrykus Ann. Rev. Plant Physiol. Plant Mol. Biol. 42:205-225 (1991).

Methods for specifically transforming dicots primarily use *Agrobacterium tumefaciens*. For example, transgenic plants reported include but are not limited to cotton (U.S. Pat. Nos. 5,004,863, 5,159,135, 5,518,908; and WO 97/43430), soybean [U.S. Pat. Nos. 5,569,834, 5,416,011; McCabe *et al.*, Bio/Technology, 6:923 (1988); and Christou *et al.*, Plant Physiol., 87:671, (1988)]; Brassica (U.S. Pat. No. 5,463,174), and peanut [Cheng *et al.*, Plant Cell Rep., 15: 653, (1996)]. Similar methods have been reported in the transformation of monocots. Transformation and

plant regeneration using these methods have been described for a number of crops including but not limited to asparagus [*Asparagus officinalis*; Bytebier *et al.*, Proc. Natl. Acad. Sci. U.S.A., 84: 5345, (1987); barley (*Hordeum vulgare*; Wan and Lemaux, Plant Physiol., 104: 37, (1994)]; maize [*Zea mays*; Rhodes, C. A., *et al.*, Science, 240: 204, (1988); Gordon-Kamm, *et al.*, Plant Cell, 2: 603, (1990); Fromm, *et al.*, Bio/Technology, 8: 833, (1990); Koziel, *et al.*, Bio/Technology, 11: 194, (1993)]; oats [*Avena sativa*; Somers, *et al.*, Bio/Technology, 10: 1589, (1992)]; orchardgrass [*Dactylis glomerata*; Horn, *et al.*, Plant Cell Rep., 7: 469, (1988); rice [*Oryza sativa*, including indica and japonica varieties, Toriyama, *et al.*, Bio/Technology, 6: 10, (1988); Zhang, *et al.*, Plant Cell Rep., 7: 379, (1988); Luo and Wu, Plant Mol. Biol. Rep., 6: 165, (1988); Zhang and Wu, Theor. Appl. Genet., 76: 835, (1988); Christou, *et al.*, Bio/Technology, 9: 957, (1991); sorghum [*Sorghum bicolor*; Casas, A. M., *et al.*, Proc. Natl. Acad. Sci. U.S.A., 90: 11212, (1993)]; sugar cane [*Saccharum* spp.; Bower and Birch, Plant J., 2: 409, (1992)]; tall fescue [*Festuca arundinacea*; Wang, Z. Y. *et al.*, Bio/Technology, 10: 691, (1992)]; turfgrass [*Agrostis palustris*; Zhong *et al.*, Plant Cell Rep., 13: 1, (1993)]; wheat [*Triticum aestivum*; Vasil *et al.*, Bio/Technology, 10: 667, (1992); Weeks T., *et al.*, Plant Physiol., 102: 1077, (1993); Becker, *et al.*, Plant, J. 5: 299, (1994)], and alfalfa [Masoud, S. A., *et al.*, Transgen. Res., 5: 313, (1996)]. It is apparent to those of skill in the art that a number of transformation methodologies can be used and modified for production of stable transgenic plants from any number of target crops of interest.

The transformed plants can be analyzed for the expression features conferred by the polynucleotides of the present invention, using methods known in the art for the analysis of transformed plants (see Example 4 of the Examples section which follows). A variety of methods are used to assess gene expression and determine if the introduced gene(s) is integrated, functioning properly, and inherited as expected. Preferably, the promoters are evaluated by determining the expression levels and the expression features of genes to which the promoters are operatively linked. A preliminary assessment of promoter function can be determined by a transient assay method using reporter genes, but a more definitive promoter assessment can be determined from the analysis of stable plants. Methods for plant analysis include but are not limited to Southern blots or northern blots, PCR-based approaches, biochemical analyses, phenotypic screening methods, field evaluations, and

immunodiagnostic assays. These methods may also be used to assess gene silencing, which is described hereinbelow.

As mentioned hereinabove, to enhance expression and/or accumulation of the molecule of interest in trichome cells and/or to facilitate purification of the molecule from trichome cells, down-regulation of at least one molecule endogenous to the plant trichomes and interfering with these processes is effected.

Trichomes are known to include a number of compounds (e.g., metabolites), which interfere with the production of molecules in these specialized cells. These metabolites include, for example polyphenols, ketones, terpenoids (e.g., monoterpenes, sesquiterpenes, diterpenes and triterpenes), mixed terpenes, phenylpropanoids and alkaloids. Other trichome components which may be preferably reduced to improve, 10 expression, accumulation and purification of the molecules of this aspect of the present invention include proteases, and PPO (see Example 5 of the Examples section). For example downregulation PPO in trichome plastids would allow the recruitment of the protein translation machinery to a novel peptide and also to increase storage space in trichome plastids. Another example is reducing enzymatic activity of the polyphenols biosynthetic pathway to thereby decrease/eliminate the production of polyphenols which make it difficult to harvest and purify proteins from trichomes (see above). Such enzymes include, but are not limited to, Phenylalanine ammonia-lyase (PAL, 15 Acc. No. M90692, M83314), Cinnamate-4-hydroxylase (CA4H, GenBank Accession No. Z70216, AI490789), 4-Coumarate:coenzyme A ligase (4CL, GenBank Accession Nos. AW034240, AF211800), chalcone and stilbene synthase (CHS, Acc. No. GenBank Accession No. X55195), Chalcone isomerase (CHI, Acc. No. GenBank Accession No. AY348871), F3H, flavanone 3-hydroxylase - naringenin 3-dioxygenase 20 (F3OH), flavanone 3-hydroxylase - naringenin 3-dioxygenase (FDR), dihydroflavonol-4-reductase (DFR, GenBank Accession No. Z18277).

Down-regulation of such trichome components may be effected by down-regulating genes which are involved in the production or accumulation of these components. For example, gene products which are involved in exudate synthesis 30 may be revealed by genome and EST mining and directed gene knock-out. Gene mining includes the identification in public databases (e.g., GenBank www.ncbi.nlm.nih.gov/Genbank/index.html) of orthologous sequences deriving from the plant of interest which share homology with known genes in the pathway using

sequence alignment software such as BLAST (www.ncbi.nlm.nih.gov/BLAST). Alternatively, trichome EST libraries may be useful for identifying genes which are involved in metabolite synthesis [see for example Lange (2000) Proc. Natl. Acad. Sci. 97:2934-2939; Gang (2001) Plant Physiology 125:539-555].

5 Once genes associated directly or indirectly with metabolite synthesis are identified, they are down-regulated either at the nucleic acid level and/or at the protein level (e.g., antibodies).

An agent capable of downregulating gene expression is a small interfering RNA (siRNA) molecule. RNA interference is a two step process. the first step, 10 which is termed as the initiation step, input dsRNA is digested into 21-23 nucleotide (nt) small interfering RNAs (siRNA), probably by the action of Dicer, a member of the RNase III family of dsRNA-specific ribonucleases, which processes (cleaves) dsRNA (introduced directly or via a transgene or a virus) in an ATP-dependent manner. Successive cleavage events degrade the RNA to 19-21 bp duplexes (siRNA), 15 each with 2-nucleotide 3' overhangs [Hutvagner and Zamore Curr. Opin. Genetics and Development 12:225-232 (2002); and Bernstein Nature 409:363-366 (2001)].

In the effector step, the siRNA duplexes bind to a nuclease complex to from 20 the RNA-induced silencing complex (RISC). An ATP-dependent unwinding of the siRNA duplex is required for activation of the RISC. The active RISC then targets the homologous transcript by base pairing interactions and cleaves the mRNA into 12 nucleotide fragments from the 3' terminus of the siRNA [Hutvagner and Zamore Curr. Opin. Genetics and Development 12:225-232 (2002); Hammond *et al.* (2001) Nat. Rev. Gen. 2:110-119 (2001); and Sharp Genes. Dev. 15:485-90 (2001)]. Although the mechanism of cleavage is still to be elucidated, research indicates that each RISC 25 contains a single siRNA and an RNase [Hutvagner and Zamore Curr. Opin. Genetics and Development 12:225-232 (2002)].

Because of the remarkable potency of RNAi, an amplification step within the RNAi pathway has been suggested. Amplification could occur by copying of the 30 input dsRNAs which would generate more siRNAs, or by replication of the siRNAs formed. Alternatively or additionally, amplification could be effected by multiple turnover events of the RISC [Hammond *et al.* Nat. Rev. Gen. 2:110-119 (2001), Sharp Genes. Dev. 15:485-90 (2001); Hutvagner and Zamore Curr. Opin. Genetics and Development 12:225-232 (2002)]. For more information on RNAi see the following

reviews Tuschl ChemBiochem. 2:239-245 (2001); Cullen Nat. Immunol. 3:597-599 (2002); and Brantl Biochem. Biophys. Act. 1575:15-25 (2002).

Synthesis of RNAi molecules suitable for use with the present invention can be effected as follows. First, the mRNA target sequence is scanned downstream of the 5' AUG start codon for AA dinucleotide sequences. Occurrence of each AA and the 3' adjacent 19 nucleotides is recorded as potential siRNA target sites. Preferably, siRNA target sites are selected from the open reading frame, as untranslated regions (UTRs) are richer in regulatory protein binding sites. UTR-binding proteins and/or translation initiation complexes may interfere with binding of the siRNA endonuclease complex [Tuschl ChemBiochem. 2:239-245]. It will be appreciated though, that siRNAs directed at untranslated regions may also be effective, as demonstrated for GAPDH wherein siRNA directed at the 5' UTR mediated about 90 % decrease in cellular GAPDH mRNA and completely abolished protein level (www.ambion.com/techlib/tm/91/912.html).

Second, potential target sites are compared to an appropriate genomic database (e.g., human, mouse, rat etc.) using any sequence alignment software, such as the BLAST software available from the NCBI server (www.ncbi.nlm.nih.gov/BLAST/). Putative target sites which exhibit significant homology to other coding sequences are filtered out.

Qualifying target sequences are selected as template for siRNA synthesis. Preferred sequences are those including low G/C content as these have proven to be more effective in mediating gene silencing as compared to those with G/C content higher than 55 %. Several target sites are preferably selected along the length of the target gene for evaluation. For better evaluation of the selected siRNAs, a negative control is preferably used in conjunction. Negative control siRNA preferably include the same nucleotide composition as the siRNAs but lack significant homology to the genome. Thus, a scrambled nucleotide sequence of the siRNA is preferably used, provided it does not display any significant homology to any other gene.

Antisense and siRNA technology has been used in selective downregulation of two tobacco trichome genes encoding different enzymes [Wang (2002); J. of Exp. Bot. 53:1891-1897; Wang (2003) Planta 216:686-691]. siRNA oligonucleotides for downregulating PPO for example, may be generated by inserting the cDNA sequence of PPO (GenBank Accession No: Z12833 for PPOA, GenBank Accession No.

Z12836 for PPOD) to an siRNA selection software such as provided by www.ambion.com.

Another agent capable of downregulating gene expression is a DNAzyme molecule capable of specifically cleaving an mRNA transcript or DNA sequence of the target. DNAzymes are single-stranded polynucleotides which are capable of cleaving both single and double stranded target sequences (Breaker, R.R. and Joyce, G. Chemistry and Biology 1995;2:655; Santoro, S.W. & Joyce, G.F. Proc. Natl. Acad. Sci. USA 1997;94:4262) A general model (the "10-23" model) for the DNAzyme has been proposed. "10-23" DNAzymes have a catalytic domain of 15 deoxyribonucleotides, flanked by two substrate-recognition domains of seven to nine deoxyribonucleotides each. This type of DNAzyme can effectively cleave its substrate RNA at purine:pyrimidine junctions (Santoro, S.W. & Joyce, G.F. Proc. Natl. Acad. Sci. USA 1999; for rev of DNAzymes see Khachigian, LM [Curr Opin Mol Ther 4:119-21 (2002)].

Examples of construction and amplification of synthetic, engineered DNAzymes recognizing single and double-stranded target cleavage sites have been disclosed in U.S. Pat. No. 6,326,174 to Joyce *et al.*

Downregulation of gene expression can also be effected by using an antisense polynucleotide capable of specifically hybridizing with an mRNA transcript encoding the target polypeptide of interest.

Design of antisense molecules which can be used to efficiently and specifically downregulate gene expression must be effected while considering two aspects important to the antisense approach. The first aspect is delivery of the oligonucleotide into the cytoplasm of the appropriate cells, while the second aspect is design of an oligonucleotide which specifically binds the designated mRNA within cells in a way which inhibits translation thereof.

In addition, algorithms for identifying those sequences with the highest predicted binding affinity for their target mRNA based on a thermodynamic cycle that accounts for the energetics of structural alterations in both the target mRNA and the oligonucleotide are also available [see, for example, Walton *et al.* Biotechnol Bioeng 65: 1-9 (1999)].

Such algorithms have been successfully used to implement an antisense approach in cells. For example, the algorithm developed by Walton *et al.* enabled

scientists to successfully design antisense oligonucleotides for rabbit beta-globin (RBG) and mouse tumor necrosis factor-alpha (TNF alpha) transcripts. The same research group has more recently reported that the antisense activity of rationally selected oligonucleotides against three model target mRNAs (human lactate dehydrogenase A and B and rat gp130) in cell culture as evaluated by a kinetic PCR technique proved effective in almost all cases, including tests against three different targets in two cell types with phosphodiester and phosphorothioate oligonucleotide chemistries.

In addition, several approaches for designing and predicting efficiency of specific oligonucleotides using an *in vitro* system were also published (Matveeva *et al.*, Nature Biotechnology 16: 1374 - 1375 (1998)).

Another agent capable of downregulating gene expression is a ribozyme molecule capable of specifically cleaving an mRNA transcript encoding the target polypeptide. Ribozymes are being increasingly used for the sequence-specific inhibition of gene expression by the cleavage of mRNAs encoding proteins of interest [Welch *et al.*, Curr Opin Biotechnol. 9:486-96 (1998)]. The possibility of designing ribozymes to cleave any specific target RNA has rendered them valuable tools in both basic research and therapeutic applications.

An additional method of regulating the expression of a gene in cells is via triplex forming oligonucleotides (TFOs). Recent studies have shown that TFOs can be designed which can recognize and bind to polypurine/polypyrimidine regions in double-stranded helical DNA in a sequence-specific manner. These recognition rules are outlined by Maher III, L. J., *et al.*, Science, 1989;245:725-730; Moser, H. E., *et al.*, Science, 1987;238:645-630; Beal, P. A., *et al.*, Science, 1992;251:1360-1363; Cooney, M., *et al.*, Science, 1988;241:456-459; and Hogan, M. E., *et al.*, EP Publication 375408. Modification of the oligonucleotides, such as the introduction of intercalators and backbone substitutions, and optimization of binding conditions (pH and cation concentration) have aided in overcoming inherent obstacles to TFO activity such as charge repulsion and instability, and it was recently shown that synthetic oligonucleotides can be targeted to specific sequences (for a recent review see Seidman and Glazer, J Clin Invest 2003;112:487-94).

In general, the triplex-forming oligonucleotide has the sequence correspondence:

		22		
oligo	3'-A	G	G	T
duplex	5'-A	G	C	T
duplex	3'-T	C	G	A

However, it has been shown that the A-AT and G-GC triplets have the greatest
5 triple helical stability (Reither and Jeltsch, BMC Biochem, 2002, Sept12, Epub). The same authors have demonstrated that TFOs designed according to the A-AT and G-GC rule do not form non-specific triplexes, indicating that the triplex formation is indeed sequence specific.

Thus, for any given sequence in the regulatory region a triplex forming
10 sequence may be devised. Triplex-forming oligonucleotides preferably are at least 15, more preferably 25, still more preferably 30 or more nucleotides in length, up to 50 or 100 bp.

Transfection of cells (for example, via cationic liposomes) with TFOs, and formation of the triple helical structure with the target DNA induces steric and
15 functional changes, blocking transcription initiation and elongation, allowing the introduction of desired sequence changes in the endogenous DNA and resulting in the specific downregulation of gene expression. Examples of such suppression of gene expression in cells treated with TFOs include knockout of episomal *supFG1* and endogenous *HPRT* genes in mammalian cells (Vasquez et al., Nucl Acids Res. 1999;27:1176-81, and Puri, et al, J Biol Chem, 2001;276:28991-98), and the sequence- and target specific down-regulation of expression of the Ets2 transcription factor, important in prostate cancer etiology (Carbone, et al, Nucl Acid Res. 2003;31:833-43), and the pro-inflammatory ICAM-1 gene (Besch et al, J Biol Chem, 2002;277:32473-79). In addition, Vuyisich and Beal have recently shown that
20 sequence specific TFOs can bind to dsRNA, inhibiting activity of dsRNA-dependent enzymes such as RNA-dependent kinases (Vuyisich and Beal, Nuc. Acids Res 2000;28:2369-74).

Additionally, TFOs designed according to the abovementioned principles can induce directed mutagenesis capable of effecting DNA repair, thus providing both
30 downregulation and upregulation of expression of endogenous genes (Seidman and Glazer, J Clin Invest 2003;112:487-94). Detailed description of the design, synthesis and administration of effective TFOs can be found in U.S. Patent Application Nos.

2003 017068 and 2003 0096980 to Froehler et al, and 2002 0128218 and 2002 0123476 to Emanuele et al, and U.S. Pat. No. 5,721,138 to Lawn.

Regardless of the methods described hereinabove, the present invention may also be effected by using mutant plants or plant variants, which do not accumulate 5 these metabolites or compounds (1, 16). Such plants can be used for expressing and/or purifying the polypeptide of interest. Alternatively, such plants can be crossed with the transgenic plants expressing the polypeptide as described hereinabove. Next generations will include plants, which both express the polypeptide of interest and produce low levels of undesired compounds.

10 Plants generated or selected according to the above is preferably capable of accumulating less than 50 % of average volume of undesired compounds in the trichome cells of the plant species.

15 The present invention also envisages a method of producing a polypeptide of interest in plant trichomes. Such polypeptides can be endogenous to the trichome or exogenous polypeptides, which can be used as pharmaceuticals (e.g., antibodies, antigens, ligands, growth factors, enzymes, structural proteins), industrial proteins and enzymes, therapeutics for veterinary use, proteins for molecular laboratories and diagnostics, nutraceuticals or cosmeceuticals.

20 The method is effected by expressing the polypeptide in the plant trichomes, as described above, and down-regulating a level of at least one molecule endogenous to the plant trichomes wherein such a molecule is capable of interfering with the expression accumulation or stability of the polypeptide of interest.

Plants which may be utilized for trichome specific expression in accordance 25 with the present invention, are preferably selected or generated capable of generating (i) a trichome size of at least 10 %, at least 20 %, at least 30 %, at least 40 %, at least 50 %, at least 80 % above average size of the plant species; (ii) leaf surface size at least 5 %, at least 10 %, at least 15 %, at least 25 %, at least 30 %, at least 40 % above average size of the plant species; and/or (iii) total leaf number at least 10 %, at least 20 %, at least 30 %, at least 40 %, at least 50 %, at least 80 % above average leaf 30 number of the plant species; (iv) trichome density on the abaxial size of the leaf at least 10 %, at least 20 %, at least 30 %, at least 40 %, at least 50 %, at least 80 % above average trichome density of the plant species; (v) trichome density on the adaxial size of the leaf at least 100 % above average trichome density of the plant

species; (vi) shoot internode length of at least 5 %, at least 10 %, at least 15 %, at least 25 %, at least 30 %, at least 40 % above average length of the plant species; (vi) trichome density above 50,000 trichomes/gr leaf tissue; and/or (vii) trichome shape different than that of the plant species.

5 Plant architecture can be designed using genetic or non-genetic approaches [Weston (1989) J. Amer. Soc. Hort. Sci. 114:492-498; Antonious (2001) J. Environ. Sci. Health B. 36(6):835-48].

A number of genes which are associated with trichome development and morphogenesis were revealed through genetic studies. These genes may regulate 10 trichome initiation, devision rate of trichome cells and/or trichome cell ploidy number. A mutation in the TTG gene (GenBank Accession Nos. TTG1 - AT5G24520 , TTG2 - AT2G37260) results in loss of leaf trichomes. Another example may be AGL16 (GenBank Accession No. NM_115583), a recently discovered MADS-box gene that is expressed in trichomes [Alvarez-Buylla (2000) Plant J. 24:457-466]. Yet another 15 example is, KIC (GenBank Accession No. AY363866), a novel Ca²⁺ binding protein with one EF-hand motif, which interacts with a microtubule motor protein and regulates trichome morphogenesis [Reddy Plant Cell. (2004) 16:185-200]. Other genes, which affect trichome size and/or distribution, include but are not limited to the UPL family of genes (e.g., UPL3, GenBank Accession No. AY265959), STICHEL 20 [GenBank Accession No. AF264023, Ilgenfritz et. al. (2003) Plant Physiol. 131:643-55], COT1 [Szymanski et.al. (1998) Genetics. 149:565-77], ZWICHEL (GenBank Accession No. AF002678, Oppenheimer et. al. (1997) Proc Natl Acad Sci U S A. 10:94:6261-6], GL1 (GenBank Accession No. AF263690), GL3 (GenBank Accession No. AT5G41315), GL2 (Acc. No. AT1G79840.1). It is conceivable that such genes 25 when over-expressed may increase trichome size and/or distribution. For example, the GL-3 homologue R gene of maize causes trichome formation when over-expressed in Arabidopsis [Schellmann (2002) EMBO J. 21:5036-5046], indicating that such a manipulation is feasible. Overexpression of heterologous genes in plants is further detailed hereinbelow.

30 Non-transgenic approaches for modifying trichome size and/or distribution include chemical or physical mutagenesis [Szymanski et. al. (1998) Plant Cell. 10:2047-62], somaclonal variation [Saied et.al. (1994) Tree Physiol. 14:17-26; Guo

et. al.(2003) Shi Yan Sheng Wu Xue Bao. 36:202-8] and induction of polyploidy [Mesaragno et. al. (1993) Plant Cell. 5:1661-1668].

Trichome density can be increased by exposure to differentiating factors (i.e., non-genetic approaches). For example, day length (16). Alternatively, physiological concentrations of ethylene have been shown to promote trichome formation [Kazama (2001) Plant Physiol. 117:375-83]. γ -radiation can be used to induce trichome formation [Negata (1999) Plant Physiol. 120:133-120].

Alternatively or additionally, trimming may be used to increase the number of leaves of the plant and as such increase the number of trichomes (see Example 7 of the Examples section).

Plants of the present invention are preferably sterile (i.e., having no viable pollen or seeds) to prevent spreading of genetic material to the surrounding environment. Sterilized plants can be selected from mutant plants produced by for example chemical mutagenesis, physical mutagenesis or by somaclonal variation. Alternatively sterilized plants can be generated by silencing of fertility genes [Siaud et. al. (2004) EMBO J. 23:1392-401; Suzuki et. al. (2004) Plant J. 37:750-61; Li et. al. (2004) Plant Cell. 16:126-43; Krishnakumar and Oppenheimer(1999) Development. 126:3079-88].

Once plants are produced in accordance with the present invention, trichome content is purified to extract the molecules expressed therein or the products thereof.

Mechanical and chemical methods of isolating trichomes and trichomes exudates and content are known in the art. Such methods include the use of solvent containing microcapillary for dissolving the exudates. Measures are taken, though, to select a solvent which does not interfere with the activity or stability of molecules thus purified. Another method for removing trichomes include the use of forceps. A more efficient method for isolating exudates is by washing the surface with an organic solvent. Again, measures are taken to select a solvent which does not interfere with the activity or stability of molecules thus purified. Trichomes may also be produced by brushing surfaces, shaking in an aqueous solution with an abrasive or freezing the tissue and then brushing [see McCaskill (1992) Planta 187:445-454; Wang (2001) Nature Biotechnology 19:371-374].

In order to facilitate collection of the trichome-produced molecule, the present inventors have devised a novel approach for large-scale collection of trichome

exudates and/or their content. This approach is simple to execute, does not require special technical skills, is cost effective and enables collection of large amounts of trichome exudates and content.

Trichome content collection according to the present invention is effected by incubating a trichome containing plant tissue in a liquid (e.g., water) such that trichome exudates and content is released into the liquid. To avoid leaching of tissue components other than trichome, liquid incubation is effected while avoiding friction between the trichome containing plant tissue and a solid surface. Thereafter, the liquid is collected, thereby harvesting the trichome exudates and content.

For example, trichome containing plant tissues, such as, shoots, leaves and flowers collected from plants can be incubated for 30-60 seconds under agitation (60 times per minute) in water or any other liquid, which allows release of trichome content and exudates, while avoiding leaching of other tissue components. Preferably avoided are non-polar solvents, such as chloroform and hexane.

The liquid is preferably supplemented with antioxidants such as citric acid, ascorbic acid and sodium bisulfite, which reduce the activity of trichome components (e.g., PPO, see Example 5 of the Examples section interfering with purification of the molecules.

Further purification of the molecules can be effected using any chemical or biochemical method known in the art depending on the chemical nature of the molecule and its intended use. Such methods include, but are not limited to, chromatography methods such as thin layer, affinity, gel filtration and ion-exchange.

Collection of trichome exudates and content can be effected manually or by employing a collection apparatus specifically designed for such a purpose.

Thus, the present invention also envisages an apparatus for mechanical harvesting of trichomes and/or trichome exudates and content (illustrated in Figure 4a), which is referred to herein as apparatus 10.

Apparatus 10 includes a collector 12 (e.g., brush, forceps, arm) which is designed and configured for collecting trichomes and/or trichome exudates and content from a trichome containing plant tissue 14. Accordingly, collector 12 includes a collecting mechanism 13 for holding plant tissue 14 and a fluid filled reservoir 16 including fluid 18 in which plant tissue 14 is agitated by collecting mechanism 13.

Reservoir 16 also serves for storing collected trichomes and/or trichome exudates and content.

As is illustrated in Figures 4b-c, to enable agitation of plant material 14 within the fluid of reservoir 16, apparatus 10 (collector 12) includes a vibrating mechanism 20 which is fitted with a motor or servo and a power unit either to collecting mechanism 13 (Figure 4b) or to reservoir 16 (Figure 4c). Apparatus 10 may also include an actuating unit 22 and a timer 24 communicating with actuating unit 22. Reservoir 16 may also include at least one liquid channel 26 and pump 28 for transferring the liquid with the trichomes out of the reservoir and into collection containers or directly to a chromatography device for further separation and molecule isolation.

Additional objects, advantages, and novel features of the present invention will become apparent to one ordinarily skilled in the art upon examination of the following examples, which are not intended to be limiting. Additionally, each of the various embodiments and aspects of the present invention as delineated hereinabove and as claimed in the claims section below finds experimental support in the following examples.

20

EXAMPLES

Reference is now made to the following examples, which together with the above descriptions, illustrate the invention in a non limiting fashion.

Generally, the nomenclature used herein and the laboratory procedures utilized in the present invention include molecular, biochemical, microbiological and recombinant DNA techniques. Such techniques are thoroughly explained in the literature. See, for example, "Molecular Cloning: A laboratory Manual" Sambrook et al., (1989); "Current Protocols in Molecular Biology" Volumes I-III Ausubel, R. M., ed. (1994); Ausubel et al., "Current Protocols in Molecular Biology", John Wiley and Sons, Baltimore, Maryland (1989); Perbal, "A Practical Guide to Molecular Cloning", John Wiley & Sons, New York (1988); Watson et al., "Recombinant DNA", Scientific American Books, New York; Birren et al. (eds) "Genome Analysis: A Laboratory Manual Series", Vols. 1-4, Cold Spring Harbor Laboratory Press, New York (1998); methodologies as set forth in U.S. Pat. Nos. 4,666,828; 4,683,202; 4,801,531;

5,192,659 and 5,272,057; "Cell Biology: A Laboratory Handbook", Volumes I-III Cellis, J. E., ed. (1994); "Current Protocols in Immunology" Volumes I-III Coligan J. E., ed. (1994); Stites et al. (eds), "Basic and Clinical Immunology" (8th Edition), Appleton & Lange, Norwalk, CT (1994); Mishell and Shiigi (eds), "Selected Methods 5 in Cellular Immunology", W. H. Freeman and Co., New York (1980); available immunoassays are extensively described in the patent and scientific literature, see, for example, U.S. Pat. Nos. 3,791,932; 3,839,153; 3,850,752; 3,850,578; 3,853,987; 3,867,517; 3,879,262; 3,901,654; 3,935,074; 3,984,533; 3,996,345; 4,034,074; 4,098,876; 4,879,219; 5,011,771 and 5,281,521; "Oligonucleotide Synthesis" Gait, M. 10 J., ed. (1984); "Nucleic Acid Hybridization" Hames, B. D., and Higgins S. J., eds. (1985); "Transcription and Translation" Hames, B. D., and Higgins S. J., Eds. (1984); "Animal Cell Culture" Freshney, R. I., ed. (1986); "Immobilized Cells and Enzymes" IRL Press, (1986); "A Practical Guide to Molecular Cloning" Perbal, B., (1984) and "Methods in Enzymology" Vol. 1-317, Academic Press; "PCR Protocols: A Guide To 15 Methods And Applications", Academic Press, San Diego, CA (1990); Marshak et al., "Strategies for Protein Purification and Characterization - A Laboratory Course Manual" CSHL Press (1996); all of which are incorporated by reference as if fully set forth herein. Other general references are provided throughout this document. The procedures therein are believed to be well known in the art and are provided for the 20 convenience of the reader. All the information contained therein is incorporated herein by reference.

EXAMPLE 1

Cloning promoter regions of trichome expressed genes and identifying trichome-active promoters

Promoters suitable for expressing proteins in trichomes were identified by sequencing the genomic DNA upstream region of various cDNAs obtained from genes expressed in leaf tissues.

MATERIALS AND METHODS

Isolation and cloning of trichome promoter sequences in a binary vector:

The NCBI database of 126,000 tomato expressed sequence tags (ESTs) (including 5,000 ESTs originated from cDNA libraries originated from the trichome tissues) and all transcribed nucleotide sequences described in literature or directly

submitted to NCBI (cDNAs) were used for the identification of trichome active promoters. Keywords representing each sequence and expression pattern thereof were collected and stored in a database.

LEADS™ software (Compugen, IL) was used for clustering and assembling
5 the tomato sequences and provided more than 20,000 clusters representing different genes. An expression profile annotative summary was designed for each cluster by pooling all keywords of each sequence represented in the cluster. Clusters were selected based on trichome EST number and percentage out of total ESTs present in each cluster. Clusters were analyzed for ORFs using Vector NTI suite (InforMax,
10 U.K.) version 6. ORFs of each gene were compared to Genbank database, using Blast (<http://www.ncbi.nlm.nih.gov/BLAST/>) and for the highest homologous ORF the position of the ATG start codon and stop codon was compared. Accordingly, most of the sequences described herein were predicted to posses the full length ORF. Clusters were classified as trichome-specific (i.e. more than 90% of ESTs in a cluster
15 were originated from trichome cDNA libraries) or trichome expressed (i.e. at least one of the ESTs in a cluster was originated from trichome cDNA libraries).

RT-PCR - To verify the levels of expression and trichome specificity Reverse Transcription following quantitative (Real-Time) PCR (RTqPCR) was performed on total RNA extracted from either leaves, trichome cells or leaves minus trichome cells.

20 mRNA levels were measured for three genes, previously predicted bioinformatically to express to high levels and specifically in trichome cells.

Trichome cells were harvested from tomato mature leaves by first freezing the leaves, just above liquid nitrogen and then brushing both sides of the leaves with paint brush, previously chilled in liquid nitrogen. Total RNA was extracted from leaves,
25 trichome cells or leaves minus trichome cells of tomato using Rneasy plant mini kit (Qiagen, Germany) using the protocol provided by the manufacturer. Reverse transcription was performed using 1.5 µg total RNA, using 300 U Super Script II Reverse Transcriptase enzyme (Invitrogen), 225ng random deoxynucleotide hexamers (Invitrogen), 500 µM dNTPs mix (Takara, Japan), 0.2 volume of x5 RT buffer
30 (Invitrogen), 0.01M DTT, 60U RNAsin (Promega), DEPC treated DDW was added up to 37.5 µl.

30

RT reactions were incubated for 50 min at 42 °C, followed by 70 °C for 15 min. cDNA was diluted 1:20 in Tris EDTA, pH=8. 5mL of the diluted cDNA was used for qPCR.

To normalize the expression level between the different tissues specific

5 primers were designed for the following housekeeping genes: Actin (SEQ ID NO: 72), GAPDH (SEQ ID NO: 73), and RPL19 (SEQ ID NO: 74). The following primers were used for qPCR: Actin F primer: CCACATGCCATTCTCCGTCT (SEQ ID NO: 77), R primer GCTTTCTTCAACGTCCCTGA (SEQ ID NO: 78); GAPDH F primer TTGTTGTGGGTGTCAACGAGA (SEQ ID NO: 79), R primer
10 ATGGCGTGGACAGTGGTCA (SEQ ID NO: 80); RPL19 F primer CACTCTGGATATGGTAAGCGTAAGG (SEQ ID NO: 81), R primer TTCTTGACTCCCTGTACTTACGA (SEQ ID NO: 82); TR2 F primer tctcttcaatttaggtacccgtttg (SEQ ID NO: 83), R primer TGAATTTCGCCGTCAATTGTCC (SEQ ID NO: 84); TR4 F primer
15 GGGTTTAGACGTATCCGAAGGTC (SEQ ID NO: 85), R primer GCTCGTTCCAATTTCAGTAGAGA (SEQ ID NO: 86); TR5 F primer TTACGTGCCAACTGAACACA (SEQ ID NO: 87), R primer CAATGCAATCAGCCCCATGC (SEQ ID NO: 88).

qPCR was performed on cDNA (5 µL), using x1 iQ™ SYBR Green super mix
20 (BioRad), forward and reverse primers 0.3 µM each, and DDW was added up to 20 µL.

qPCR reaction was performed in iCycler real-time PCR machine (BioRad) 95 °C for 3min, 40 times of 95 °C for 15 sec and 1min at 60 °C, followed by 95 °C for 15 sec, 60 °C for 60 sec, and 70 times of 60 °C for 10 sec +0.5 °C increase in each cycle.

25 The levels of expression (Qty) measured in the qPCR were calculated using the efficiency (E) of the amplification reaction and the corresponding C.T. (the cycle at which the samples crossed the threshold) Qty=E^{C.T.}. This calculation method assumes that the efficiencies of the reactions of the GOI (gene of interest) and of the housekeeping genes are similar. In general the efficiencies of the reactions were 100%
30 +/- 5%. .

Results are summarized in Figures 6a-d.

Figures 6a-d show that all three selected genes (i.e. TR2, TR4, and TR5) were expressed at high leves, up to 21 times higher than the housekeeping genes, in

trichome cells. In all cases expression was higher in trichomes compared to leaves, and in *L. hirsutum* compared to *L. esculentum* plants. Hence the promoter sequences, upstream to the gene sequences, were cloned from *L. hirsutum* gDNA.

In order to clone these promoter sequences and 5' untranslated region (5' UTR) upstream of the ATG starting codon, total genomic DNA was extracted from plant leaf tissues of 4 week old plants of the following species: cultivated tomato (*Lycopersicon esculentum*, var 870), wild tomato species (*Lycopersicon hirsutum*, var LA 1777 and *Lycopersicon pennellii*, var LA 716), tobacco (*Nicotiana tabaccum*, var NN) or cotton (*Gossypium hirsutum* var Acala 23). DNA extraction was effected using DNA extraction kit (Dneasy plant mini kit, Qiagen, Germany). Inverse PCR (IPCR), DNA digestion, self-ligation, and PCR reaction were performed on genomic DNA, following a well established protocol ([http://www.pmc1.unimelb.edu.au/core facilities/manual/mb390.asp](http://www.pmc1.unimelb.edu.au/core_facilities/manual/mb390.asp)) with the following modifications. To avoid mistakes in the IPCR, first the genomic sequence of the 5' sequence of a relevant cDNA (i.e. including introns) was identified to produce Genomic Island (GI). The desired region from the genomic DNA was PCR-amplified using direct oligonucleotide primers designed based on the cDNA cluster sequence, as was predicted by the Leads software (CompuGen, IL). PCR reaction was performed in a DNA thermal cycler, using common PCR conditions (for example: 92 °C/3 min followed by 31 cycles × [94 °C/30 sec; 56 °C/30 sec; 72 °C/3 min] followed by 72 °C/10 min). PCR products were purified using PCR purification kit (Qiagen) and sequencing of the amplified PCR products was performed, using ABI 377 sequencer (Amersham Biosciences Inc.).

Primer sequences of each plant and the resultant GI sequence (i.e. the genomic sequence which was amplified using the primers) are listed in Table 1, below.

32
Table 1

<i>ID/Plant</i>	<i>Forward primer/SEQ ID NO:</i>	<i>Reverse primer/SEQ ID NO:</i>	<i>Product size/SEQ ID NO:</i>
TR2 (<i>L. hirsutum</i>)	atggaagtaactttgttgtatagtac/SEQ ID NO: 1	GCCAGTGATCACCATAAGGA G/SEQ ID NO: 2	376/SEQ ID NO: 3
TR4 (<i>L. hirsutum</i>)	Ttcttggtcttcaatgttgg/SEQ ID NO: 4	TTTGTAAATGTCATTGGGAGGT C/SEQ ID NO:5	410 bp of 5' prime region out of about 3500 bp of amplified PCR product SEQ ID NO: 6; Note - 3500 bp were amplified by PCR, out of which only 5' prime 410 bp were sequenced
TR5 (<i>L. hirsutum</i>)	Gggtaatattcatttgattttcc/SEQ ID NO: 6	AACCTGCTTACATGTTCAA G/ SEQ ID NO: 7	431 bp/ SEQ ID NO: 9

To increase amplification efficiency as needed a different amplification technique [UP-PCR (20)] was employed. Briefly, UP-PCR technique was used in order to amplify unknown upstream region of a known cluster sequence. Generally, the procedure involved four oligonucleotide primers: two sequence specific primers (SPs, external and internal) (listed below), both having the same orientation of 3' end towards the unknown, yet desired, 5' region of the gene, and two universal walking primers (WP28 and sWP). Reaction mixtures were generated as follows: sample mixture (SM) - genomic DNA of appropriate plant (tomato or cotton) species (30-40ng), WP28 primers (20 pmol), and DDW was added to a final volume of 10 μ l; Polymerase mixture (PM) – dNTPs (Roche, Switzerland) (10mM each), Expand Long Template Enzyme mix (Roche, Switzerland) (1U), 10 X buffer supplied with the enzyme and DDW was added to a final volume of 8 μ l. SM was placed in a thermocycler (Biometra, USA), where it was subjected to an amplification program of 1 minute at 90 °C, held (pause) at 80 °C until PM was added, 30 seconds at 15 °C, 10

minutes at 25 °C, 3 minutes at 68 °C, held at 90 °C until the external SP (2 µl of 10 µM concentration) was added. The process was followed by external PCR reaction of 30 seconds at 92 °C, 10 seconds at 94 °C, 30 seconds at 65.5 °C, 3 minutes at 68 °C, for 30 cycles followed by final extension of 10 minutes at 68 °C.

External PCR products (diluted 5000 – 25000 fold) were used as template and subjected to amplification using specific internal sWP and SP (30 pmol each) primers, 1U Ex Taq (Takara), in 50µl reaction volume. Internal PCR reactions were subjected to an amplification program of 2 minutes at 92 °C, followed by 30 seconds at 94 °C, 30 seconds at 58 °C, and 3 minutes at 72 °C for 30 cycles and a final extension of 10 minutes at 72 °C. IPCR/Up-PCR products were purified (PCR Purification Kit, Qiagen, Germany) and sequenced (ABI 377 sequencer, Amersham Biosciences Inc). Table 2, below, lists primers and products of IPCR/Up-PCR reactions.

15

Table 2

<i>ID/Plant</i>	<i>Amplification method</i>	<i>External primers</i>		<i>Internal primers</i>		<i>Product SEQ ID NO:</i>
		<i>sWP28/SEQ ID NO:</i>	<i>SP (external)/SEQ ID NO:</i>	<i>sWP/SEQ ID NO:</i>	<i>SP (internal)/SEQ ID NO:</i>	
TR2/ <i>L. hirsutum</i>	UP-PCR	TTTTTTTT TTGTTTGT TGTGGGGG TGT/10	GGAAGTTT AAGTAGTG GGCTTG/11	TTTTG TTTGTT GTGGG/ 12	GTGGGCTT GGTGGTAG ATTC/13	14
TR4/ <i>L. hirsutum</i>	UP-PCR	TTTTTTTT TTGTTTGT TGTGGGGG TGT/10	GTTGAGTC CACGAGCA GACAC/15	TTTTG TTTGTT GTGGG/ 12	CGAGCAGAC ACTGTCAGA GG/16	17
TR5/ <i>L. hirsutum</i>	UP-PCR	TTTTTTTT TTGTTTGT TGTGGGGG TGT/10	ATTCACAA GGTTGTGG ATGAGG/18	TTTTG TTTGTT GTGGG/ 12	GATGAGGT GTTTGGGT GCAC/19	20

For cloning the putative promoters and 5' UTRs, an additional PCR amplification was effected using a new set of primers (below) which included 8-12 bp extensions having one restriction site (*Hind*III, *Sal*II, *Xba*I, *Bam*HI, or *Sma*I) on the 5'

prime end thereof. For each promoter, restriction sites that do not exist in the promoter sequence were selected. Moreover, the restriction sites in the primer sequences were design such that the resultant PCR products were cloned into the binary vector pPI in the right orientation, upstream of the GUS reporter gene.

5 The plasmid pPI was constructed by inserting a synthetic poly-(A) signal sequence, originating from pGL3 basic plasmid vector (Promega, Acc No U47295; bp 4658-4811) into the *Hind*III restriction site of the binary vector pBI101.3 (Clontech, Acc. No. U12640).

10 Table 3, below lists the restriction enzymes containing primers, and the SEQ ID NO: of the resultant PCR products. Restriction sites within each primer are indicated by bold letters.

Table 3

<i>ID/Plant</i>	<i>Forward primer - Restriction enz./SEQ ID NO:</i>	<i>Reverse primer - Restriction enz./SEQ ID NO:</i>	<i>Product SEQ ID NO:</i>
TR2 <i>(L. hirsutum)</i>	(<i>Hind</i> III): 5'- AATTAAAGCTTGTGTCGCTC AGCCCCTACTC -3'/21	(<i>Sall</i>): 5'- AAATTGTCGACATCTCAAC TTGTTGCACTGAATTG -3'/22	23
TR4 <i>(L. hirsutum)</i>	(<i>Sall</i>): 5'- CCTAGTCGACGGTGTAAA TGGTGGGTTGG -3'/24	(<i>Bam</i> HI): 5'- TTGGATCCGAGCAGACACT GTCAGAGG -3'/25	26
TR5 <i>(L. hirsutum)</i>	(<i>Hind</i> III): 5'- TTCCAAGCTTGACCTGCTC TGATACCAATTG -3'/27	(<i>Bam</i> HI): 5'- CCGGATCCTCGTAAGGAGT TTGTAATATG -3'/28	29

15 PCR products were purified (PCR Purification Kit, Qiagen, Germany) and digested with the restriction sites according to the primers used (Roche, Switzerland). The digested PCR products were re-purified and cloned into the binary vector pPI, which was digested with the same restriction enzymes. PCR product and the linearized plasmid vector were ligated using T4 DNA ligase enzyme (Roche, 20 Switzerland).

EXAMPLE 2***Cloning of trichome active promoter sequences***

Several genomic sequences were already described and validated in the literature as trichome-specific promoters. In most cases promoter validation was effected in tobacco plants. However, none of these sequences were analyzed in tomato plants. Hence there is no way to predict which of these promoters will be active in tomato.

MATERIALS, METHODS AND RESULTS

A previously described tobacco promoter, (8) was isolated from genomic DNA (gDNA) of *Nicotiana tabaccum*, var Samsun NN using two sets of overlapping primers: 1. Forward- 5'- AAAT**T**CTAGACTACCATCGCTAGTAATCGTG -3' (SEQ ID NO: 30) and Reverse- 5'- GTTGAAGAACTGCATCCC**G**GGGAGG -3' (SEQ ID NO: 31) to provide the sequence product set forth in SEQ ID NO: 32 (, TR25-2, SEQ ID NO: 67); 2. Forward- 5'- AAAT**T**CTAGATAAGTTGATAAAGCTAATTCTC -3' (SEQ ID NO: 33) and Reverse- 5'- TTT**C**CCCGGGACCTGGAGGCAATC -3' (SEQ ID NO: 34) to provide the sequence product set forth in SEQ ID NO: 35 (TR25-3, SEQ ID NO: 68).

Primers sequences included additional restriction sites XbaI (Forward primers) and SmaI (Reverse primers), indicated in bold letters, to facilitate further cloning.

Each PCR product was digested with XbaI and SmaI (Roche) and introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with the same restriction enzymes.

A cotton promoter, previously described in (7), was isolated from gDNA of *Gossypium hirsutum*, var. Acala 23, and *Gossypium barbadense* var. Pima 15 using the primers: Forward- 5'- TATAAGCTTAAAGTTAAATCCTATTGTAGTG -3' (SEQ ID NO: 36) and Reverse- 5'- CGGATCCATTAAATCACAAAGAAAAAC -3' (SEQ ID NO: 37) to provide a genomic amplified sequence of Acala as set forth in SEQ ID NO: 38 (27A) and a genomic amplified sequence of Pima as set forth in SEQ ID NO: 39 (27P).

Primer sequences included additional restriction sites HindIII (Forward primer) and BamHI (Reverse primer), indicated in bold letters, to facilitate further cloning. PCR products were digested with HindIII and BamHI (Roche) and

36

introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with the same restriction enzymes.

EXAMPLE 3

5 Cloning of tomato PPO promoters and signal peptide

Tomato polyphenol oxidase (PPO, GenBank Accession No: Z12833 for PPOA, GenBank Accession No. Z12836 for PPOD) is the major protein in type VI trichomes of tomato (5). Hence it was expected that the promoter region upstream the PPO gene will direct the expression of foreign genes to the trichome cells. PPO is encoded by closely related, seven members, gene family. A previous publication identified which of the gene family members are preferably expressed in the trichome cells (5).

The genomic sequence of the PPO gene family was published. Still, in most cases, promoter activity was not tested for the sequences upstream of the genes.

15 MATERIALS, METHODS AND RESULTS

The promoter sequence of PPOA and PPOD was cloned from wild tomato (*Lycopersicon pennellii*) and cultivated tomato (*Lycopersicon esculentum*), respectively. Cloning of the putative promoter region of PPOA was effected by amplifying the genomic sequence upstream of the coding region, using the primers:

20 Forward- 5'- AAAATTGGGATCTAGAAGGTGAGG -3' (SEQ ID NO: 40) and Reverse- 5'- CTGGATCCTATTGCTAGCTTGGATGAAG -3' (SEQ ID NO: 41). The resultant genomic DNA amplified thereby is set forth in SEQ ID NO: 42 (T8). Primer sequences include additional restriction sites XbaI (Forward primer) and BamHI (Reverse primer), indicated in bold, to facilitate further cloning.

25 The resultant PCR product was digested with XbaI and BamHI (Roche) and introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with the same restriction enzymes.

Cloning of the putative promoter region of PPOD was performed by amplifying the genomic sequence upstream of the coding region, using the primers:

30 Forward- 5'- ATGGAAAAGCTTATGGACAGACTAAACAC -3' (SEQ ID NO: 43) and Reverse- 5'- CTGGATCCTGTTGCTAGCTTTGAATGAAA -3' (SEQ ID NO: 44). The resultant genomic DNA amplified thereby is set forth in SEQ ID NO: 45 (T11).

Primer sequences included additional restriction sites HindIII (Forward primer) and BamHI (Reverse primer), indicated in bold, to facilitate further cloning.

PCR product was digested with HindIII and BamHI (Roche) and introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with the same 5 restriction enzymes.

Vast accumulation of PPO in trichomes is largely governed by protein import and storage within the thylakoid lumen of plastids, such as chloroplasts and leucoplasts (5, 12, 28). Protein import into the lumen is directed by a signal peptide on the amino terminal of the immature polypeptide of PPO. The immature 10 immature polypeptide is imported first into the plastid stroma where the primary part of the signal peptide is cleaved. Later on, the second part of the signal peptide is cleaved, while the polypeptide is crossing the thylakoid membrane and the mature polypeptide is entering into the thylakoid lumen (28).

Hence to facilitate the accumulation of foreign proteins in trichomes, protein 15 import into plastids might be crucial.

To test protein import, the native signal peptide of either PPOA or PPOD was amplified together with the putative promoter of each gene. Amplified products were cloned into pPI and fused, in frame, to the GUS reporter gene. Amplification of PPOA promoter together with only the initial part of the signal peptide, which directs 20 protein to the stroma was done using the following primers: Forward- 5'-AAAATTGGGATCTAGAAGGTGAGG -3' (SEQ ID NO: 46, XbaI restriction site is indicated in bold) and Reverse- 5'- ACATGAAACTTGAATGCTTG-3' (SEQ ID NO: 47). The genomic amplified sequence of PPOA is set forth in SEQ ID NO: 48.

25 PCR product was digested with XbaI (Roche) and introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with XbaI and SmaI restriction enzymes.

Amplification of PPOD promoter and signal peptide which directs protein to the stroma was effected using the following primers: Forward- 5'- 30 ATGGAAAAGCTTATGGACAGACTAAAACAC -3' (SEQ ID NO: 49) and Reverse- 5'-TTCCCGGGACATGAAACTTGAATGCTTG -3' (SEQ ID NO: 50). Genomic amplified sequence of PPOD is set forth in SEQ ID NO: 51.

Primer sequences included additional restriction sites HindIII (Forward primer) and SmaI (Reverse primer) to facilitate further cloning.

PCR product was digested with HindIII and SmaI (Roche) and introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with the same 5 restriction enzymes.

Amplification of PPOD promoter and signal peptide which directs protein to the lumen was done using the following primers: Forward- 5'-ATGGAAAAGCTTATGGACAGACTAAACAC -3' (SEQ ID NO: 52) and Reverse- 5'-AACCCGGGAGCCGATGCAGCTAATGG -3' (SEQ ID NO: 53). The 10 resultant genomic amplified sequence of PPOD is set forth in SEQ ID NO: 54.

Primer sequences included additional restriction sites HindIII (Forward primer) and SmaI (Reverse primer), indicated in bold, to facilitate further cloning.

PCR product was digested with HindIII and SmaI (Roche) and introduced via ligation, using T4 DNA ligase (Roche), into pPI plasmid, digested with the same 15 restriction enzymes.

EXAMPLE 4

Expression of therapeutic proteins in trichome cells

MATERIALS AND METHODS

20 The potential of trichomes to accumulate human therapeutic proteins can be estimated by expressing human interferon β gene or human growth hormone gene in trichome cells.

MATERIALS AND METHODS

25 ***Cloning of human interferon β into a binary vector*** - The gene for human interferon β (INFB, GenBank Accession No. NM_002176) was amplified from human genomic DNA using the following primers: Forward: 5'-GGGATGAGCTACAACTTGCTTGGAT-3' (SEQ ID NO: 55) and Reverse: 5'-CTAGGAGCTCTCAGTTCGGAG-3' (SEQ ID NO: 56, a SacI restriction site on the primer is indicated in bold). The resultant sequence of the INFB gene is set forth 30 in SEQ ID NO: 57.

Analysis of the INFB sequence revealed a codon usage that is similar to the codon usage of the tomato (data not shown).

The PCR product (SEQ ID NO: 57) was digested using *SacI* restriction endonuclease (Roche) and cloned into pPI binary vector digested with *SmaI* and *SacI*, hence replacing the GUS gene. The newly formed binary plasmid was designated pINFB. Sequence analysis of the INFB gene in pINFB revealed that the INFB sequence was cloned in the right orientation. Trichome promoters together or without a plastid transit peptides (summarized in Table 4 below) were further cloned upstream of the INFB gene in pINFB.

5 *Cloning of human growth hormone into a binary vector* - The mature polypeptide of the Human-growth hormone gene (HGH, GenBank Accession No: V00519) was produced synthetically using GeneArt service (<http://www.geneart.de/>). The sequence was adjusted according to the tomato codon usage, while avoiding, as much as possible, high GC content and low complexity of DNA sequences. An ATG was added as a first codon to the mature polypeptide enabling sufficient translation. The restriction sites of *SmaI* and *SacI* were added to the gene at the 5' prime end and 10 3' prime end, respectively. The sequence of the HGH gene is set forth in SEQ ID NO: 58.

15 The gene clone was provided in a PCR script plasmid vector. The gene was digested out of the plasmid using *SmaI* and *SacI* restriction endonucleas (Roche) and cloned into pPI binary vector, replacing the GUS gene. The newly formed binary plasmid was named pHGH. Sequence analysis revealed that the inserted HGH gene in pHGH was cloned in the right orientation. Trichome promoters with or without a 20 plastid transit peptides (summarized in Table 4, below) were further cloned upstream of the HGH gene in pHGH.

25 *Agrobacterium transformation of binary plasmids expressing heterologous genes* - *Agrobacterium tumefaciens* (strains LBA4404) competent cells were transformed with 0.5 µl binary plasmid by electroporation, using a MicroPulser electroporator (Biorad, USA), 0.2 cm cuvettes (Biorad, USA) and EC-2 electroporation program (Biorad, USA). Cells were incubated in LB medium at 28 °C for 3 hours and plated on LB-agar plates supplemented with 50 mg/L kanamycin 30 (Sigma, USA) and 250 mg/L streptomycin. Plates were incubated at 28 °C for 48 hours until *Agrobacterium* colonies grew. These colonies were subsequently used for tobacco or tomato plant transformation.

Plant transformation and cultivation – Table 4, below, summarizes the constructs which were introduced into tomato plants.

Table 4

Promoter	species	Transit peptide	gene	binary
35S	CaMV	No	GUS	pPI
TR2	<i>L. hirsutum</i>	No	GUS	pPI
TR4	<i>L. hirsutum</i>	No	GUS	pPI
TR5	<i>L. hirsutum</i>	No	GUS	pPI
TR8	<i>L. pennellii</i>	No	GUS	pPI
TR8	<i>L. pennellii</i>	Stroma	GUS	pPI
TR11	<i>L. esculentum</i>	No	GUS	pPI
TR11	<i>L. esculentum</i>	Stroma	GUS	pPI
TR11	<i>L. esculentum</i>	Lumen	GUS	pPI
TR25-2	<i>N. tabaccum</i>	No	GUS	pPI
TR25-3	<i>N. tabaccum</i>	No	GUS	pPI
TR27-A	<i>G. hirsutum</i>	No	GUS	pPI
TR27-P	<i>G. barbadense</i>	No	GUS	pPI
TR2	<i>L. hirsutum</i>	No	INFβ	pINFβ
TR4	<i>L. hirsutum</i>	No	INFβ	pINFβ
TR5	<i>L. hirsutum</i>	No	INFβ	pINFβ
TR8	<i>L. pennellii</i>	No	INFβ	pINFβ
TR8	<i>L. pennellii</i>	Stroma	INFβ	pINFβ
TR11	<i>L. esculentum</i>	No	INFβ	pINFβ
TR11	<i>L. esculentum</i>	Stroma	INFβ	pINFβ
TR11	<i>L. esculentum</i>	Lumen	INFβ	pINFβ
TR25-2	<i>N. tabaccum</i>	No	INFβ	pINFβ
TR25-3	<i>N. tabaccum</i>	No	INFβ	pINFβ
TR27-A	<i>G. hirsutum</i>	No	INFβ	pINFβ
TR27-P	<i>G. barbadense</i>	No	INFβ	pINFβ
TR2	<i>L. hirsutum</i>	No	HGH	pHGH
TR4	<i>L. hirsutum</i>	No	HGH	pHGH
TR5	<i>L. hirsutum</i>	No	HGH	pHGH
TR8	<i>L. pennellii</i>	No	HGH	pHGH
TR8	<i>L. pennellii</i>	Stroma	HGH	pHGH
TR11	<i>L. esculentum</i>	No	HGH	pHGH
TR11	<i>L. esculentum</i>	Stroma	HGH	pHGH
TR11	<i>L. esculentum</i>	Lumen	HGH	pHGH

TR25-2	<i>N. tabaccum</i>	No	HGH	pHGH
TR25-3	<i>N. tabaccum</i>	No	HGH	pHGH
TR27-A	<i>G. hirsutum</i>	No	HGH	pHGH
TR27-P	<i>G. barbadense</i>	No	HGH	pHGH

Tomato transformation - Tomato transformation was carried out according to Fillati *et al.* (19). Briefly, *Lycopersicon esculentum* cv. Micro-Tom cotyledons were used for *Agrobacterium* based plant transformation. The Micro-Tom seeds were 5 surface sterilized for 10 min in a 3 % sodium hypochlorite solution. The seeds were washed with DDW three times and soaked for three hours in fresh DDW, then plated into 0.5 L container with Nitsch medium, containing MS salts, 3 % sucrose, Nitsch vitamins and 0.8 % plant agar (Duchefa, Netherland).. The PH was adjusted to 5.8 prior to autoclaving for 20 min at 121 °C. 50 seeds where plated on 0.5 L sterilized 10 container containing the germination medium and left at 25 °C in culture room, 16/8 hrs light/dark cycles, under light intensity of (150 $\mu\text{Em}^{-2}\text{s}$).. Seedlings where grown for 8 days.

Agrobacterium tumefaciens strain LBA4404 carrying an intact vir region which can mediate the introduction of the T-DNA from the bacteria into plants. The 15 binary vector plasmids, originated from pPI, were introduced into the strain LBA 4404 as described above.

For co cultivation, a single colony from freshly streak LB plate supplemented with 300 $\mu\text{g}/\text{ml}$ streptomycin and 50 $\mu\text{g}/\text{ml}$ kanamycin (Sigma) was used to inoculate 5 ml LB overnight shaking at 28 °C. The inoculated 5 ml where added to 45ml LB 20 supplemented with 300 $\mu\text{g}/\text{ml}$ streptomycin and 50 $\mu\text{g}/\text{ml}$ kanamycin to additional overnight at the same conditions. The overnight culture where centrifuged for 3060 rpm for 10 minutes and rinsed with 50 ml MS medium.

2.5 ml of fine tobacco suspension culture where plated on petri dishes (100x25mm) containing 50 ml of KCMS murashige minimal organics medium 25 supplemented with 0.2 $\mu\text{g}/\text{ml}$ 2.4D, kinetin 0.1 $\mu\text{g}/\text{ml}$, thiamine hydrochloride 0.8 $\mu\text{g}/\text{ml}$, potassium acid phosphate 200 $\mu\text{g}/\text{ml}$, biotin 0.5 $\mu\text{g}/\text{ml}$, folic acid 0.5 $\mu\text{g}/\text{ml}$, casein hidrolysat 800 $\mu\text{g}/\text{ml}$ and plant agar 0.8%, PH 5.8 (Duchefa, Netherland)..

Whatmann paper filter no.1 (Whatmann) was autoclaved and placed on top the of the feeder plates. Any air bubbles and remaining tobacco suspension media were

excluded. The plates were incubated for 24 hours under low light conditions ($10 \mu\text{Em}^{-2}\text{s}$).

8 days old cotyledons were cut at both ends on MS medium and plated for 24 hours on the tobacco suspension plates in the same conditions.

5 The cotyledons were immersed in 5 ml of the rinsed *Agrobacterium* cells diluted in 50 ml MS medium in sterile Petri dish. The concentration of the bacteria was $\sim 5 \times 10^8$ cfu /ml. Following 10 minutes the cotyledons were blotted carefully to remove any excess of bacterial suspension. 20 cotyledons were placed on the feeder plates for 48 hr for co-incubation with the bacteria under the same conditions.

10 Cotyledons were then transferred to 2 Zeatin Ribozide (ZR) regeneration medium (Duchefa, Netherland), containing 400 $\mu\text{g}/\text{ml}$ carbenicillin or 150 Ticarcillin/ potassium clavulanate to inhibit growth of *Agrobacterium* and Kanamycin 100 $\mu\text{g}/\text{ml}$ to select for transformed tomato cells.

15 The cotyledons were transferred to fresh regeneration media after 1 month supplemented with 1 ZR 200 carbenicillin and 100 $\mu\text{g}/\text{ml}$ Kanamycin.

20 Shoots where excised when they were approximately 1 cm long and transferred to 0.5 L containers supplemented with rooting media containing MS medium 50 $\mu\text{g}/\text{ml}$ kanamycin, 100 $\mu\text{g}/\text{ml}$ carbenicillin disodium, 2 $\mu\text{g}/\text{ml}$ IBA (Duchefa, Netherland). After approximately ten days the rooted explants were transferred into soil, under 100% humidity. The humidity was reduced gradually for 24 hours. After 24 hr the plants were transferred to the greenhouse.

Testing expression of Foreign Proteins in Trichomes:

25 **A. GUS staining** - Gus staining of tomato and tobacco plants was effected as previously described (15). Briefly: Leaves were fixed in 90 % ice cold acetone for 15 - 20 minutes (on ice), followed by removal of acetone and a double tissue rinsing with the Working Solution [25mM Sodium Phosphate (Sigma, USA) buffer pH=7, Ferricyanide (Sigma, USA) 1.25mM, Ferrocyanide (Sigma, USA) 1.25mM, Triton X-100 (Sigma, USA) 0.25%, EDTA(BioLab) 0.25mM] for 15-20 minutes. Rinse solution was removed, replaced with Staining solution [Working solution with 5-bromo-4-chloro-3-indolyl- β -D-glucuronic acid (X-GlcA, Duchefa) solubilized in N,N-Dimethylformamide (BioLab) 1.5mg/ml and Dithiothreitol (DTT, Bio Lab) 100mM] and incubated in the dark (tubes wrapped with aluminum foil) over night at 37 °C. Distaining was effected by sinking the plant tissue in 70 % ethanol and heating

at 50 °C for ~ 2 hours. The distaining step was repeated until the plant tissue became transparent except the blue stained regions. Distained plants were stored in 70 % ethanol (BioLab) at room temperature.

5 **ELISA** - Human protein detection in plant tissues was effected using Human Interferon β ELISA Kit (R & D Systems) and Ultra-sensitive Human Growth Hormone ELISA Kit (Diagnostic Systems Laboratories, Inc), according to manufacturer instructions.

10 **Western Blot** - Briefly, proteins extracted from the leaves were resolved on 12 % Tris-HCl Criterion gel (Bio-Rad Laboratories, Inc.) and transferred by electroporation onto PVDF membranes using the Bio-Rad Criterion Precast Gel System (100 V constant voltage at 4 °C, 1.5 hours). Pre-stained SDS-PAGE standards (Bio-Rad Laboratories, Inc.) were used as molecular weight markers. Primary antibodies were diluted 1:500 and the secondary antibody-HRP conjugate was diluted 1:15,000. The following anti- recombinant protein polyclonal antibodies 15 were used: goat (Santa Cruz) or sheep (Biosource) polyclonal anti human IFN- β , goat polyclonal anti human GH (Santa Cruz), rabbit polyclonal anti *E.coli* β -glucuronidase (Molecular Probes). ECL and related reagents were obtained from Amersham Biosciences.

20 **Bradford protein quantification test** - Protein quantification was effected according to Bradford method [Bradford M., Analytical biochemistry, 72: 248 (1976)] using Bio-Rad Laboratories, Inc. reagent.

RESULTS

25 GUS staining was performed on leaves of T1 tomato and tobacco plants transfected with a binary vector including putative trichome specific promoters upstream of a GUS gene. Control plants included wild-type tomato and tomato transfected with GUS under the constitutive CaMV S35 promoter (Figures 1b-c). As is shown in Figures 1d-h under the regulation of TR2, TR5, TR11, TR25 and TR27 promoters GUS was expressed in a trichome-gland specific manner (Figures 1d-h). A light blue color was also found in the stalk cell immediately attached to the gland and 30 in non glandular tissues. Results for tomato T1 generation are summarized in Table 6, below.

44
Table 6

Promoter	Signal peptide	No of Independent T1 plants	Avg grade	Range
35S	No	9	2.1	0-5
TR2	No	2	1.5	0-3
TR5	No	6	0.2	0-1
TR11	No	3	0	0
TR11	Stroma	4	0	0
TR11	Lumen	4	0.5	0-1
TR25-2	No	8	0	0
TR25-3	No	18	0.1	0-1
TR27-P	No	5	0.4	0-2

The results presented in Table 6 above indicate that TR2 and TR27-P are most effective in facilitating expression of heterologous genes in the tomato trichome.

5 These results are of special significance since TR2 has never been described before as a trichome active promoter. Furthermore, the addition of a luman directing signal peptide seems to have facilitated expression in the trichome.

EXAMPLE 5

10 *Novel methods for mechanical harvesting of proteins from trichomes*

MATERIALS AND METHODS

Harvesting of Trichome proteins:

Four harvesting protocols were attempted to optimize protein purification.

15 *Protocol 1* - Total trichome protein harvesting was performed by wiping fully expanded leaves with cotton swabs moistened with a solution of 200 mM Dithiothreitol (DTT, BioLab). Approximately 5-7 leaves were wiped per swab, with a total of about 1500 leaves. The crude trichome extract was squeezed from the swab using a syringe, and centrifuged (15000 x g for 15 minutes at 4 °C). To adsorb phenolic compounds, the supernatant (approximately 70 ml) was treated with 20 Polyvinylpolypyrrolidone (PVPP; Sigma, USA) for 10 minutes on a gentle stirrer followed by centrifugation (15000g for 20 minutes) to remove the PVPP. Prior to application, PVPP was boiled in 10 % HCl for 10 minutes, washed extensively with DDW, air-dried for storage and soaked for at least 3 hours in 200 mM DTT solution

at 4 °C (ratios were 8-12 g PVPP / 250ml of 200mM DTT, 1-1.5g / 60-85ml extract). Extracts were concentrated in two steps of ammonium sulfate precipitation (20 %, 75 % of saturation). Solid ammonium sulfate (0%→ 20%: 114g/1L, 20%→ 75%: 382g/1L) was added to the extract until the desired concentration was reached while 5 constant stirring. Thereafter the solution was held on ice for about 1 hour with occasional stirring. The precipitate was collected by centrifugation (15000 X g for 20 minutes at 4 °C) and dissolved in 50 mM Tris-HCl, pH 7 and 30mM NaCl.

Protocol 2 - Trichome cells and exudates were harvested directly into DDW which contained chemicals with antioxidant activity (i.e., citric acid, ascorbic acid or 10 sodium bisulfite). Tomato leaves and shoots were soaked in a container. Gentle shaking of the tissue in the liquid medium caused the explosion of the glandular trichome cells, type VI and VII (see Figure 1a) and released the trichome exudates into the media. To eliminate the loss of significant amounts of liquid, tissues were lifted out of the medium and partially dried by shaking it vigorously, letting drops fall 15 back to the container. Trichome yield was measured by inspecting the treated leaves. Trichome harvesting efficiency was calculated as the percentage of broken and exploded trichomes, out of total trichomes on a given leaf area.

Protocol 3 - Trichome cells and exudates were harvested directly into a liquid media. Tomato leaves or shoots were put in a container, filled with tap water. The 20 container was closed and centrifuged at 20 rpm for 5 minutes. To eliminate the loss of significant amounts of liquid, tissues were lifted out of the medium and partially dried by shaking it vigorously, letting drops fall back to the container. Trichome yield was measured by inspecting the treated leaves. Trichome harvesting efficiency was calculated by percentage of broken and exploded trichomes, out of total trichomes on 25 a given leaf area.

Protocol 4 - Trichome cells and exudates were harvested directly into a liquid media. Tomato leaves or shoots were put in a container, filled with tap water. The container was closed and water was poured on top of the leaves using pump which circulated the water in the container. Trichome yield was measured by inspecting the 30 treated leaves. Trichome harvesting efficiency was calculated by percentage of broken and exploded trichomes, out of total trichomes on a given leaf area.

RESULTS

Protein harvest, purification, yield – Protocol 1 was effected on a commercial tomato variety, grown in a commercial greenhouse, which was designed and used for tomato fruit harvest. Tomato (*L. esculentum* var 591) plants were grown for 3 months. Plant architecture was designed by leaving two main shoots for each plant. Using Bradford analysis it was possible to calculate total protein yield. Harvesting about 1,100 leaves yielded 16 mg total protein (0.1 mg/ml). Total protein yield was resolved on Nu-PAGE Novex Bis-Tris gel, 12 % (Invitrogen) and protein bands were visualized by coomassie staining (Figure 2). A band harboring of an estimated size of 60 kDa was predicted to be the mature PPO protein according to previous reports (12, 29). PPO is estimated to count for 60 % of total proteins in the trichome.

It will be appreciated that although protocol 1 for protein harvesting is highly effective in collecting most of the trichome exudates, it is labor intensive and slow. The other described methods (2, 3, and 4) were tested to replace the mechanical harvesting step in method 1.

Tomato shoots (1 m long) were dipped in a container and were subjected to trichome mechanical harvesting using method 2, 3 or 4.

Results, showing the efficiency of each method are presented in the Table 7, below.

20

Table 7

<i>Method</i>	<i>Number of Shoots</i>	<i>Average Efficiency</i>	<i>Range Efficiency</i>
2	3	98	96-100
3	3	11.3	8-14
4	3	42	38-46

These results suggest that protocol 2 is the most effective for protein harvesting from trichomes.

High activity of PPO in trichome may affect protein harvesting and purification from trichomes. Thus, identification of chemicals which are strong antioxidants, non toxic, cheap and not affecting protein stability is mostly desired.

Three compounds were tested: citric acid, ascorbic acid and sodium bisulfite. Different concentrations of chemicals were used to identify the chemical with the highest antioxidant activity. Tomato (*L. esculentum*) young leaves (about 100mg)

47

were grinded at 4 °C in 200 μ l of 10 mM Tris-HCl pH=8 buffer containing appropriate concentration of antioxidant (0.01 - 2.0 %w/v), incubated for about 2 hrs at 4°C and centrifuged at 14,000 rpm for 3 min in order to separate leaves-debris and liquid fraction. Decreasing of PPO activity was inspected by eliminating the production of brown color of the supernatant as a result of the oxidation of polyphenols. Table 8 below summarizes the minimal concentration that enables a decrease of 95% of PPO activity.

Table 8

<i>chemical</i>	<i>Efficient concentration</i>
Citric acid	1.0%
Ascorbic acid	1.0%*
Sodium bisulfite	0.05%

10 * After overnight incubation at 4 °C all the samples undergone browning regardless of the ascorbic acid concentration.

Figure 3 shows browning of medium as a result of PPO activity, under different concentrations of Sodium bisulfite.

15 A buffer without any antioxidant was used as a negative control. DDT was used as a positive control at the concentration of 200 mM, which is known to eliminate PPO activity when harvested from trichome cells (12, 29).

EXAMPLE 6

An automated machine for trichome cells and trichome exudate harvesting

A machine was designed, to enable the automation and up-scaling of protein harvesting according to protocol 2. The purpose of the machine is to harvest trichome cells and exudates from full-grown, 3 month old tomato plants (about 1 meter long and 80 cm in diameter). The machine is built of 2 main parts, (i) 2 m high and 1 m in diameter of cylinder shaped container (either made from stainless-steel, glass or a plastic); and (ii) a 2.5 m arm which operates using a 2 speed engine.

The machine has four steps of operation:

1. The plant is tight to the arm manually. The arm is introducing the plant into the container, half full with the liquid medium.
2. The arm is slowly moving up and down (engine is operating on a slow speed) the plant in the liquid medium such that trichome cells and exudates are released into the medium, without damaging other tissues of the plant.

3. The arm is moving the plant just above of the liquid medium.
4. The arm is vigorously shaking the plant up and down (engine is operating on a high speed). Doing that most of the liquid, which remained attached to the plant tissues, is released and drops falls back to the container. The semi-dried
5. plant is removed and a new plant is tight to the arm.

EXAMPLE 7

Determining plant architecture for optimizing trichome production

Typically, the architecture of cultivated tomato plants is designed via breeding

10 to provide the highest fruit yields, in a given space, in a given time. Moreover hand labor is routinely being practiced to optimize plant architecture for that purpose. For trichome optimized expression, plant architecture needs re-design to optimize trichome production, in a given greenhouse space in a given time. Two approaches for increasing protein yield were employed essentially, increasing the number of

15 trichomes in leaves; and increasing the number of leaves on the plant.

MATERIALS METHODS AND RESULTS

Increasing number of trichomes on leaves - Over 300 tomato cultivars were screened for trichome density. Leaves of 4 weeks old plants were inspected, and average trichome density was measured. Best performing cultivars were grown and

20 trichome density was tested again on mature, 14 week old plants. Trichome density was compared to previously measured density of several tomato lines (1). The seven best performing cultivars were grown in the next season and trichome density was measured again, to check the heredity of trichome density of two generations. Table 9, below summerizes trichome density of the best performing cultivars.

25

Table 9

<i>Var</i>	<i>No. of Plants 1st generation</i>	<i>Avg. trichome number of 1st generation</i>	<i>No of Plants 2nd generation</i>	<i>Avg. trichome number of 2nd generation</i>
309_2	3	6	3	5.3
305	3	4.7	3	5.3
249_1	3	4.3	3	5.3
247	3	3.7	3	6
273_1	3	5.4	3	4.7

294	3	4.5	3	3.2
289	3	4.2	3	3

(Note – the number in the above table represent only the best performing cultivars, out of 300 tested).

Folding a single leaflet and inspecting the edge of the folded leaflet was performed in order to count trichome cells. Trichome number is all trichomes found 5 on the edge of the leaflet under X 120 magnification using binocular microscope (Optika, Italy). Previous publication has calculated for *Lycopersicon hirsutum*, var *Glabratum* (Cultivar No 273_1 in this experiment) over 100,000 trichomes per 1 gram of leaf. Assuming trichome density in this experiment remains the same, best performing *L. esculentum* cultivars (No 309_2, 305, 249_1, 247, 294, 289), identified 10 here, have the trichome number in the same order. Each leaflet was inspected 3-5 times and an average number was calculated for the leaflet. Three different leaflet from three different plants were inspected for each cultivar in each generation.

Four cultivars were identified with the highest density of type VI (Figure 1a) 15 glandular trichomes on the upper part of the leaves. Among the five best performing cultivars, one belongs to *Lycopersicon hirsutum*, var *Glabratum*. (273_1) and the rest for cultivated tomato (No 309_2, 305, 249_1, 247,). Interestingly, the *Lycopersicon esculentum* species cultivars exhibited up to 20 fold more coverage of type VI trichomes compared to other cultivars within this species (not shown). Overall best 20 cultivated cultivars possessed the same density of type VI trichomes, compared to the wild species *Lycopersicon hirsutum*, var *glabratum*, which is recognized as the highest trichome density in all *Lycopersicon* genus (1, 16).

Approach B - Increasing number of leaves in a plant - Tomato plant architecture was designed manually. 35 days old plants were planted in a greenhouse. Different mechanical treatments were applied to shape plant architecture during plant 25 growth. To avoid the collapse of the plant bush, shoots were hanged from the greenhouse ceiling using plastic strings. The three best performing treatments for plant architecture, aiming to increase trichome yield by increasing the number of leaves produced are presented hereinbelow:

1. Plant shoot number is not limited, plant height is limited to 1m, flowers 30 were cut-off before fruit set.

50

2. Plant shoot number is not limited, plant height is limited to 2m, flowers were cut-off before fruit set.

3. Plant shoot number is not limited, leading apical meristem was cut (i.e. breaking apical dominance) when reached 0.5m, flowers were cut-off before fruit set.

5 4. Plant shoot number was limited to two. Flowers and fruits remained untouched (A control treatment, usually applied for greenhouse tomatoes, grown for fruit set).

10 Three different indeterminant (i.e. the greenhouse type) tomato cultivars (namely 678, 1312, and 2545) previously identified as having high trichome density, were grown in the greenhouse for two months. The four above treatments were applied to five plants from each cultivar. Leaf number of each plant was calculated. Table 10, below summarizes the leaf number of tomato plants growing under different mechanical plant design .

Table 10

Var	treat	N Rows	Mean(No)	Std Err(No)
678	1	5	94.2	7.61
678	2	5	87.4	5.22
678	3	5	71	5.03
678	4	5	27.8	2.75
1312	1	5	125	24.56
1312	2	5	148.4	13.12
1312	3	5	123	10.22
1312	4	5	54.6	2.84
2545	1	5	95.6	23.89
2545	2	5	81.4	3.91
2545	3	5	81.8	12.92
2545	4	5	34.4	4.48

15

As is evident from Table 10, above and Figures 5a-c, a significant increase [at 0.05 level for cultivar 2545 (Figure 5c) and 0.01 level for cultivars 678 and 1312 (Figures 5a-b, respectively)] was observed in response to treatments number 1, 2, 3 compared to control (treatment No 4). Overall a 50 % increase in leaf number (2.25 to 20 3.39) was observed over the control. Trichome density and PPO enzyme activity in trichomes were measured in each plant to verify that the increase in leaf number is not correlated with a decrease in trichome or protein production. No significant change (at 0.05 level) was observed for either trichome number or protein accumulation, following growth in leaf number (data not shown).

It is appreciated that certain features of the invention, which are, for clarity, described in the context of separate embodiments, may also be provided in combination in a single embodiment. Conversely, various features of the invention, which are, for brevity, described in the context of a single embodiment, may also be 5 provided separately or in any suitable subcombination.

Although the invention has been described in conjunction with specific embodiments thereof, it is evident that many alternatives, modifications and variations will be apparent to those skilled in the art. Accordingly, it is intended to embrace all 10 such alternatives, modifications and variations that fall within the spirit and broad scope of the appended claims. All publications, patents and patent applications mentioned in this specification are herein incorporated in their entirety by reference into the specification, to the same extent as if each individual publication, patent or patent application was specifically and individually indicated to be incorporated 15 herein by reference. In addition, citation or identification of any reference in this application shall not be construed as an admission that such reference is available as prior art to the present invention.

REFERENCES CITED

(Additional references are cited in the text)

1. Antonious GF. 2001. Production and quantification of methyl ketones in wild tomato accessions. *J Environ Sci Health B.* ;36(6):835-48.
2. Phillips MA, Croteau RB. 1999. Resin-based defenses in conifers. *Trends Plant Sci.* 4(5):184-190.
3. Wagner GJ. 1991. Secreting glandular trichomes: more than just hairs. *Plant Physiol.* 96, 675-679.
4. Kennedy GG. 2003. Tomato, pests, parasitoids, and predators: tritrophic interactions involving the genus *Lycopersicon*. *Annu Rev Entomol.* 48:51-72.
5. Thipyapong P, Joel DM, Steffens JC. 1997. Differential Expression and Turnover of the Tomato Polyphenol Oxidase Gene Family during Vegetative and Reproductive Development. *Plant Physiol.* 113(3):707-718.
6. Wang E, Wang R, DeParasis J, Loughrin JH, Gan S, Wagner GJ. 2001. Suppression of a P450 hydroxylase gene in plant trichome glands enhances natural-product-based aphid resistance. *Nat Biotechnol.* 19(4):371-4.
7. Liu HC, Creech RG, Jenkins JN, Ma DP. 2000. Cloning and promoter analysis of the cotton lipid transfer protein gene Ltp3(1). *Biochim Biophys Acta.* 24;1487(1):106-11.
8. Wang E, Gan S, Wagner GJ. 2002. Isolation and characterization of the CYP71D16 trichome-specific promoter from *Nicotiana tabacum* L. *J Exp Bot.* 53(376):1891-7.
9. Hsu CY, Creech RG, Jenkins JN, Ma DP. 1999. Analysis of promoter activity of cotton lipid transfer protein gene LTP6 in transgenic tobacco plants. *Plant Sci* 143, 63-70.
10. Wagner GJ. 1999. Tobacco surface chemistry. In *Tobacco production chemistry and technology*. (eds Davis, D.L. & Nielsen, M.T.) 292-303 (Blackwell Science, Malden, MA; 1999).
11. Kowalsky SP. 1989. Insect resistance in potato: purification and characterization of a polyphenol oxidase from the type A glandular trichomes of *Solanum berthaultii* Hawkes. PhD thesis Cornell University, Ithaca, NY.

12. Kowalsky SP, Eannetta NT, Hirzel AT, Steffens JC. 1992. Purification and characterization of a polyphenol oxidase from the type A glandular trichomes of *Solanum berthaultii*. *Plant Physiol.* 100, 677-684.
13. Duke SO *et al.* 1999. Sequestration of phytotoxins by plants: Implications for biosynthetic production. In *Biologically active natural products: agrochemicals* (eds Cutler HG & Cutler SJ) 127-136 (CRC press Boca Raton, FL; 1999).
14. Kesley RG, Reynolds GW, Rodriguez E. 1984. The chemistry of biologically active constituents secreted and stored in plant glandular trichomes. In *Biologically and chemistry of plant trichomes* (eds Rodriguez E, Healey PL, Mehta I) 187-241 (Plenum press New York; 1984).
15. Jefferson RA, Kavanagh TA, Bevan MW. 1987. GUS fusions: beta-glucuronidase as a sensitive and versatile gene fusion marker in higher plants. *EMBO J.* 6(13): 3901-7.
16. Weston PA, Johnson DA, Burton HT, Snyder JC. 1989. Trichome secretion composition, trichome densities, and spider mite resistance to ten accessions of *Lycopersicon hirsutum*. *J. Amer. Soc. Hort. Sci.* 114(3): 492-498.
17. Clough SJ, Bent AF. 1998. Floral dip: a simplified method for Agrobacterium-mediated transformation of *Arabidopsis thaliana*. *Plant J.* 16(6): 735-43.
18. Desfeux C, Clough SJ, Bent AF. 2000. Female reproductive tissues are the primary target of Agrobacterium-mediated transformation by the *Arabidopsis* floral-dip method. *Plant Physiol.* 123(3): 895-904.
19. Fillatti (1987) Efficient transfer of a glyphosate tolerance gene into tomato using a binary *Agrobacterium tumefaciens* vector. *Bio/Technology* 5 726-730.
20. Dominguez O., C. Lopez-Larrea. 1994. Gene walking by unpredictably primed PCR. *Nucleic Acids Research*. 22:3247-3248.
21. Hareven D, Gutfinger T, Parnis A, Eshed Y, Lifschitz E. 1996. The making of a compound leaf: genetic manipulation of leaf architecture in tomato. *Cell*. 84:735-44.
22. Hu Y, Xie Q, Chua NH. 2003. The *Arabidopsis* auxin-inducible gene ARGOS controls lateral organ size. *Plant Cell*. 15:1951-61.

23. Kim JH, Choi D, Kende H. 2003. The AtGRF family of putative transcription factors is involved in leaf and cotyledon growth in *Arabidopsis*. *Plant J.* 36:94-104.
24. Smith HM, Hake S. 2003. The interaction of two homeobox genes, BREVIPEDICELLUS and PENNYWISE, regulates internode patterning in the *Arabidopsis* inflorescence. *Plant Cell.* Aug. 15:1717-27.
25. Sorefan K, Booker J, Haurogne K, Goussot M, Bainbridge K, Foo E, Chatfield S, Ward S, Beveridge C, Rameau C, Leyser O. 2003. MAX4 and RMS1 are orthologous dioxygenase-like genes that regulate shoot branching in *Arabidopsis* and pea. *Genes Dev.* 17:1469-74.
26. Rose AB, Last RL. 1997. Introns act post-transcriptionally to increase expression of the *Arabidopsis thaliana* tryptophan pathway gene PAT1. *Plant J.* 11:455-64.
27. Morello L, Bardini M, Sala F, Breviaro D. 2002. A long leader intron of the Ostub16 rice beta-tubulin gene is required for high-level gene expression and can autonomously promote transcription both in vivo and in vitro. *Plant J.* 29:33-44.
28. Sommer A, Ne'eman E, Steffens J.C, Mayer A.M, Harel E. 1994. Import, targeting and processing of a plant Polyphenol oxidase. *Plant Physiol.* 105:1301-1311.
29. Yu H, Kowalsky S.P, and Steffens J.C. 1992. Comparison of Polyphenol Oxidase expression in glandular trichomes of *Solanum* and *Lycopersicon* species. *Plant Physiol.* 100, 1885-1890.
30. Pérez-Estrada L.B, Cano-Santana Z and Oyama K. 2000. Variation in leaf trichomes of *Wigandia urens*: environmental factors and physiological consequences. *Tree Physiol.*, 20:629-632

WHAT IS CLAIMED IS:

1. An isolated polynucleotide comprising a nucleic acid sequence at least 80 % identical to SEQ ID NO: 23, 26 or 29, wherein said nucleic acid sequence is capable of regulating expression of at least one polynucleotide sequence operably linked thereto in trichomes.
2. A nucleic acid construct comprising the isolated polynucleotide of claim 1.
3. The nucleic acid construct of claim 2, wherein the nucleic acid construct further comprising at least one heterologous polynucleotide operably linked to the isolated polynucleotide.
4. The nucleic acid construct of claim 3, wherein the nucleic acid construct further comprises, a nucleic acid sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a subcellular compartment of a trichome.
5. The nucleic acid construct of claim 4, wherein said nucleic acid sequence is selected from the group consisting of SEQ ID NOs: 59, 61, 63, 65 and 67.
6. The nucleic acid construct of claim 4, wherein said subcellular compartment of a trichome is a leucoplast.
7. A transgenic cell comprising the nucleic acid construct of claim 2.
8. A transgenic plant comprising the nucleic acid construct of claim 2.
9. An isolated polynucleotide comprising a nucleic acid sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a subcellular compartment of a trichome, wherein said peptide is encoded by the polynucleotide sequence set forth in SEQ ID NO: SEQ ID NOs: 59, 61, 63, 65 and 67.

10. A nucleic acid construct comprising the isolated polynucleotide of claim 9.

11. The nucleic acid construct of claim 10, further comprising an expressible polynucleotide sequence translationally fused to said nucleic acid sequence encoding said peptide.

12. A method of producing a polypeptide of interest in plant trichomes, the method comprising:

- (a) expressing the polypeptide of interest in the plant trichomes; and
- (b) down-regulating a level of at least one molecule endogenous to the plant trichomes, said at least one molecule being capable of interfering with expression, accumulation or stability of the polypeptide of interest.

13. The method of claim 12, wherein said expressing the polypeptide of interest in the plant trichomes is effected by introducing into the plant trichomes a nucleic acid sequence encoding the polypeptide of interest positioned under a transcriptional control of a promoter functional in the plant trichomes.

14. The method of claim 13, wherein said promoter is as set forth in SEQ ID NO: 23, 26, 29, 35, 38, 39, 42, 48, 50 or 51.

15. The method of claim 13, wherein said nucleic acid sequence encoding the polypeptide of interest further encodes a peptide capable of directing transport of the polypeptide of interest fused thereto into a subcellular compartment of the plant trichomes.

16. The method of claim 15, wherein said subcellular compartment of the plant trichomes is a leucoplast.

17. The method of claim 12, wherein said at least one molecule endogenous to the plant trichomes is an enzyme or a metabolite.

18. The method of claim 17, wherein said metabolite is selected from the group consisting of polyphenols, ketones, terpenoids, phenylpropanoids and alkaloids.

19. The method of claim 17, wherein said enzyme is PPO.

20. The method of claim 17, wherein step (b) is effected by gene silencing.

21. A method of producing a molecule of interest in plant trichomes, the method comprising:

- (a) expressing a polypeptide capable of directly or indirectly increasing a level of the molecule of interest in the plant trichomes; and
- (b) down-regulating a level of at least one molecule endogenous to the plant trichomes, said at least one molecule being capable of interfering with accumulation or stability of the molecule of interest, thereby producing the molecule in the plant trichomes.

22. The method of claim 21, wherein said polypeptide is endogenously expressed in the plant trichomes.

23. The method of claim 21, wherein said expressing said polypeptide in the plant trichomes is effected by introducing into the plant trichomes a nucleic acid sequence encoding said polypeptide positioned under a transcriptional control of a promoter functional in the plant trichomes.

24. The method of claim 23, wherein said promoter is as set forth in SEQ ID NO: 23, 26, 29, 35, 38, 39, 42, 48, 50 or 51.

25. The method of claim 23, said nucleic acid sequence encoding the polypeptide of interest further encodes a peptide capable of directing transport of said polypeptide fused thereto into a subcellular compartment of the plant trichomes.

26. The method of claim 25, wherein said subcellular compartment of the plant trichomes is a leucoplast.

27. The method of claim 21, wherein said at least one molecule is an enzyme or a metabolite.

28. The method of claim 27, wherein said metabolite is selected from the group consisting of polyphenols, ketones, terpenoids, phenylpropanoids and alkaloids.

29. The method of claim 27, wherein said enzyme is PPO.

30. The method of claim 27, wherein step (b) is effected by gene silencing.

31. A plant genetically modified to express a molecule of interest in trichomes, wherein said plant is further modified or selected capable of accumulating less than 50 % of average volume of undesired compounds in trichome cells of said plant species.

32. The plant of claim 31, wherein at least a portion of cells of the plant are genetically modified to include an expression construct including a polynucleotide sequence of a trichome specific promoter.

33. The plant of claim 32, wherein said expression construct further includes an additional polynucleotide sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a subcellular compartment of said trichome, whereas said additional polynucleotide is translationally fused to said polynucleotide sequence.

34. The plant of claim 31, wherein at least a portion of cells of the plant are genetically modified to include an expression construct including a first polynucleotide sequence encoding said polypeptide translationally fused to a second polynucleotide sequence encoding a peptide capable of directing transport of a polypeptide fused thereto into a trichome.

35. The plant of claim 31, wherein said expression or accumulation is in a subcellular compartment of trichomes.

36. The plant of claim 35 wherein said subcellular compartment is a leucoplast.

37. The plant of claim 32, wherein said trichome specific promoter is set forth by SEQ ID NO: 23, 26 or 29.

38. The plant of claim 32, wherein said trichome specific promoter is set forth by SEQ ID NO: 23, 26, 29, 35, 38, 39, 42 or 45.

39. The plant of claim 33, wherein said additional polynucleotide sequence is set forth by SEQ ID NO: 59, 61, 63, 65 or 67.

40. The plant of claim 31, wherein said molecule of interest is not a reporter polypeptide.

41. The plant of claim 31, wherein said plant is modified or selected capable of generating a trichome density above 50,000 trichomes/gr leaf tissue.

42. The plant of claim 31, wherein said plant is modified or selected capable of generating a trichome size of 50 % above average size of said plant species.

43. The plant of claim 31, wherein said plant is modified or selected capable of generating leaf surface size at least 25 % above average size of said plant species.

44. The plant of claim 31, wherein said plant is modified or selected capable of generating total leaf number at least 50 % above average leaf number of said plant species.

45. The plant of claim 31, wherein said plant is sterile.

46. The plant of claim 31, wherein said plant is further genetically modified capable of secreting said exogenous polypeptide from trichome cells.

60

47. A method of harvesting trichomes and/or exudates and/or content thereof, the method comprising:

- (a) incubating a trichome-containing plant tissue in a liquid such that trichome exudates and content is released into said liquid, wherein incubating is effected while avoiding friction of said trichome-containing plant tissue with a solid surface; and
- (b) collecting said liquid, to thereby harvest the trichome exudates and content.

48. The method of claim 47, wherein said liquid includes an antioxidant.

49. The method of claim 48, wherein said antioxidant is selected from the group consisting of citric acid, ascorbic acid and sodium bisulfite

50. The method of claim 49, wherein said liquid is water.

51. The method of claim 47, wherein said trichome-containing plant tissue is selected from the group consisting of a shoot, a flower and a leaf.

52. An apparatus for mechanical harvesting of trichome exudates and content, the apparatus comprising a mechanism designed and configured for mechanically agitating a trichome-containing plant tissue in a fluid and collecting said fluid to containing the trichome exudates or content.

1/17

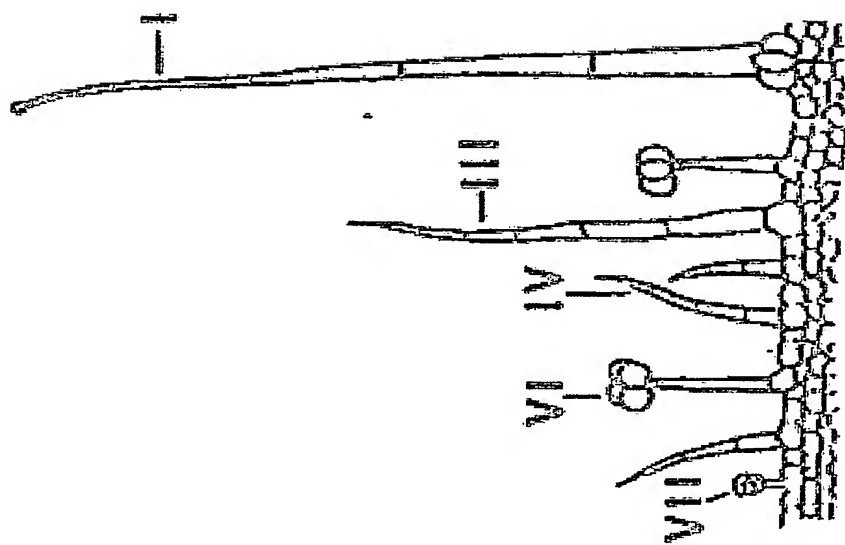
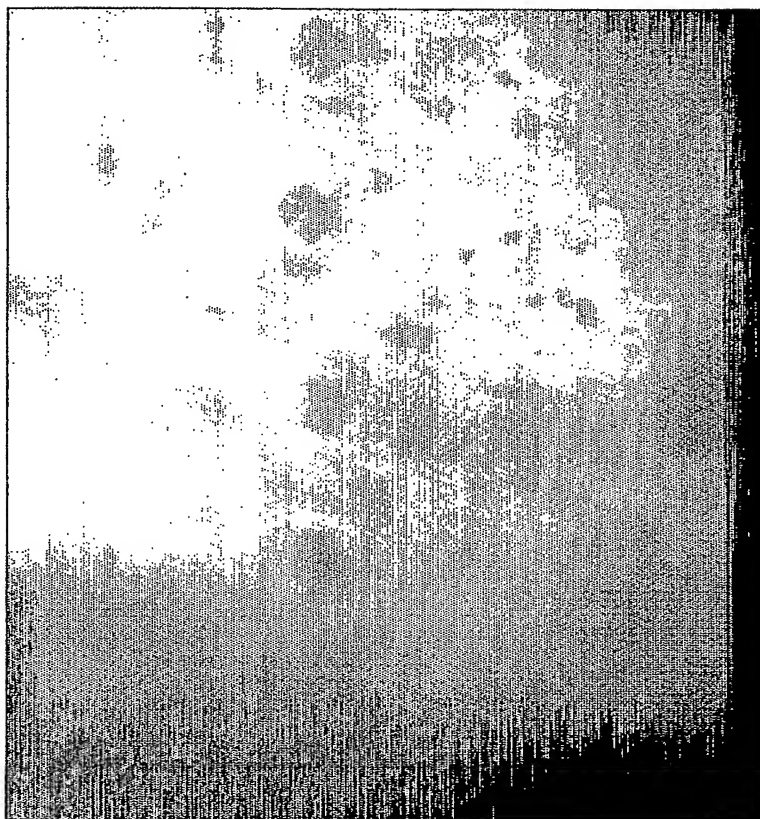


Fig. 1a

2/17



W.T.

Fig. 1b

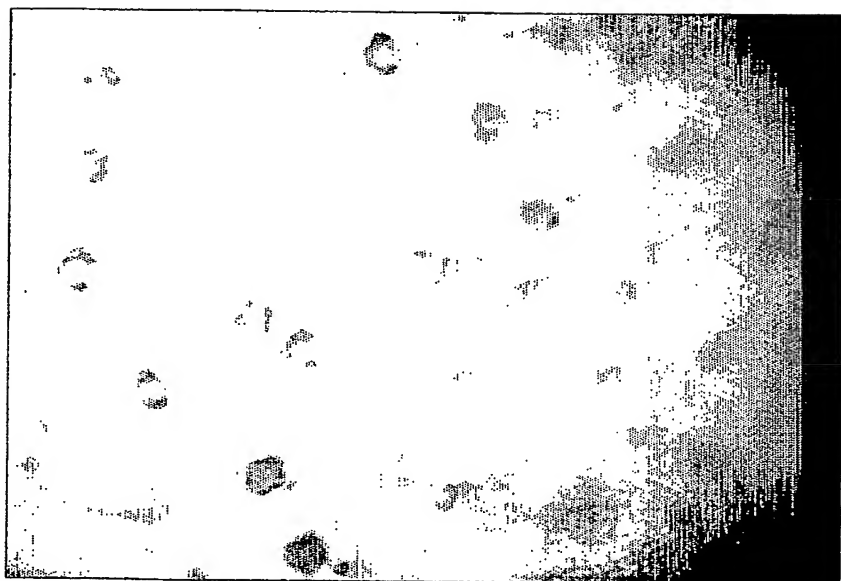
3/17



35s

Fig. 1c

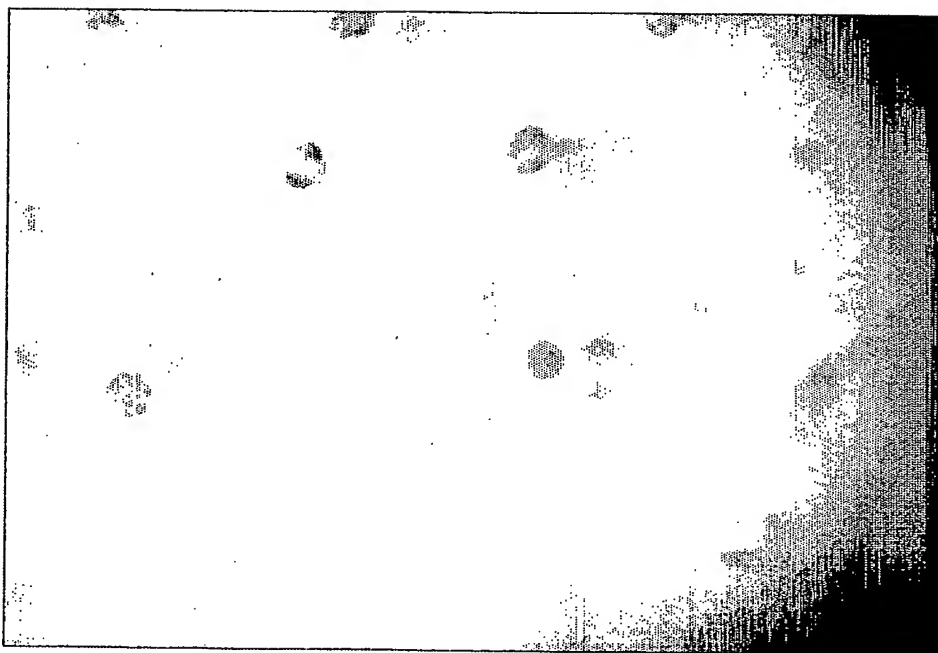
4/17



TR2H

Fig. 1d

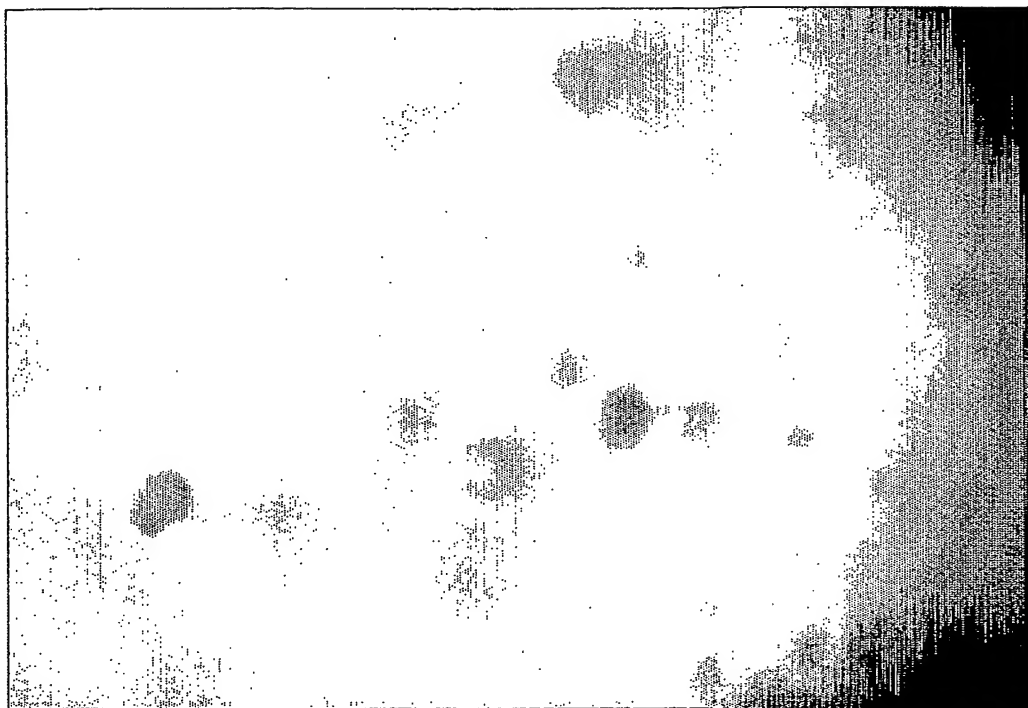
5/17



TR5H

Fig. 1e

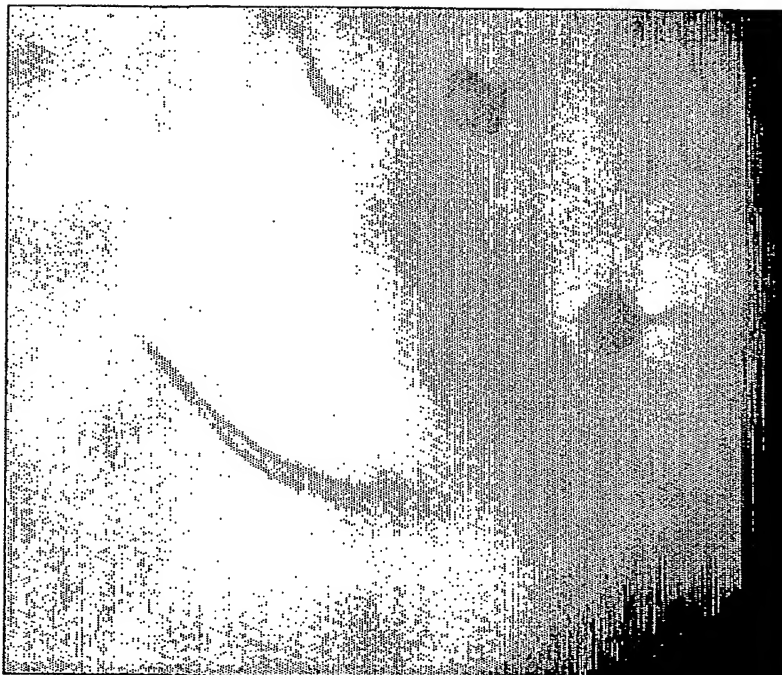
6/17



TR11E

Fig. 1f

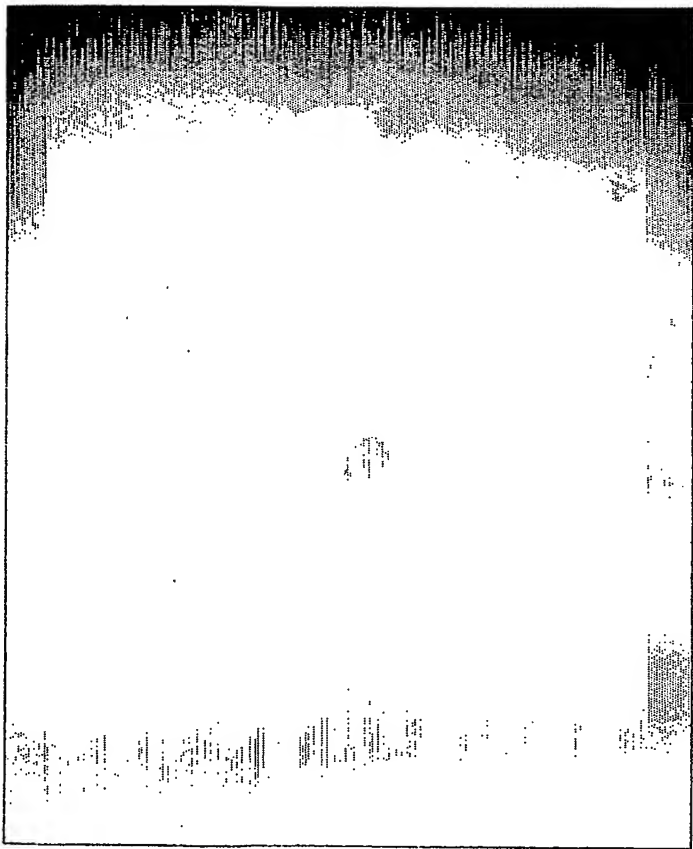
7/17



TR25

Fig. 1g

8/17



TR27P

Fig. 1h

9/17

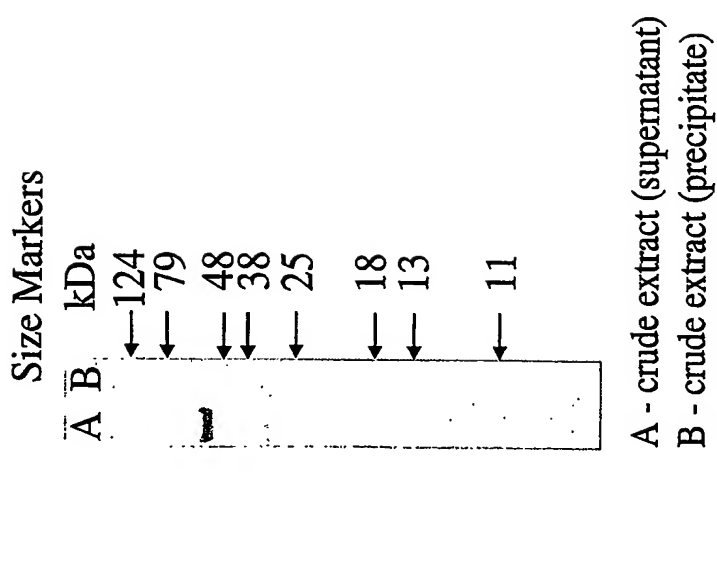


Fig. 2

10/17

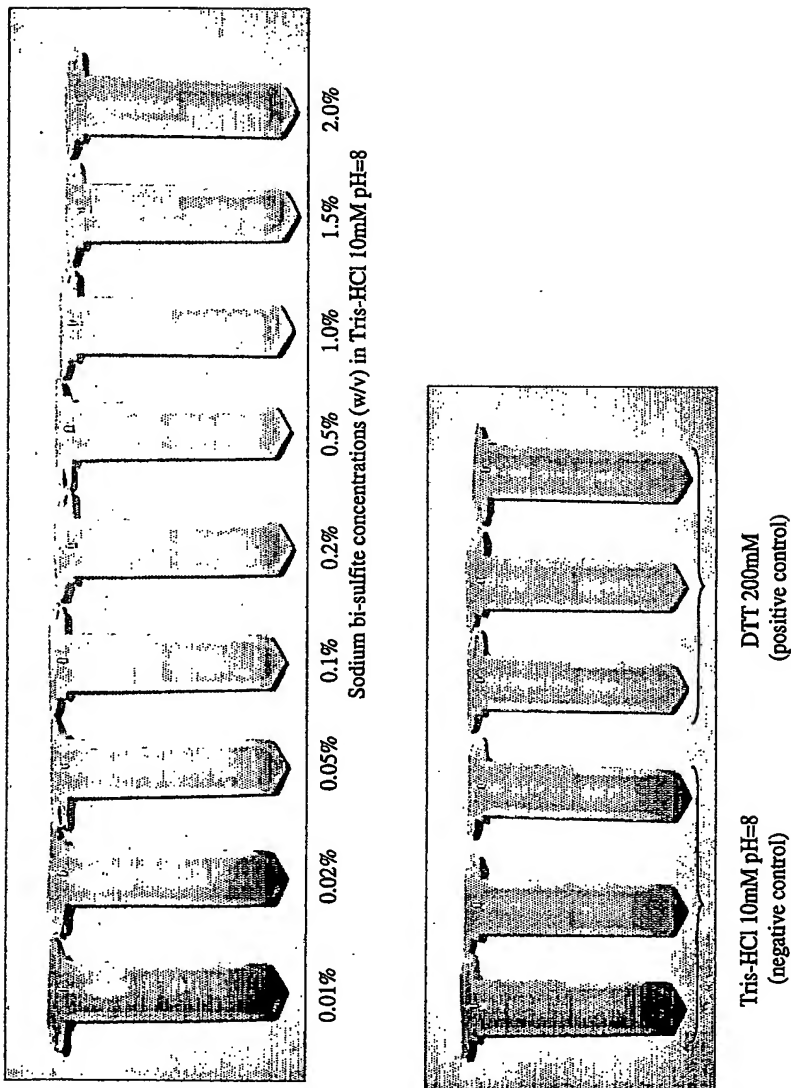


Fig. 3

11/17

10

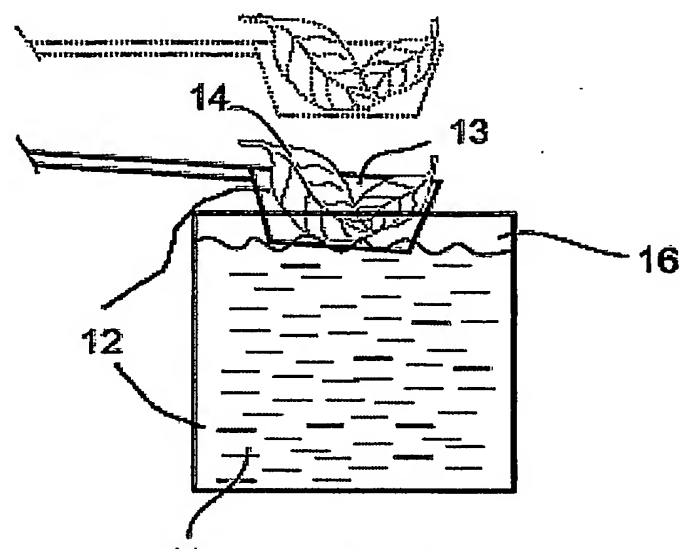


Fig. 4a

10

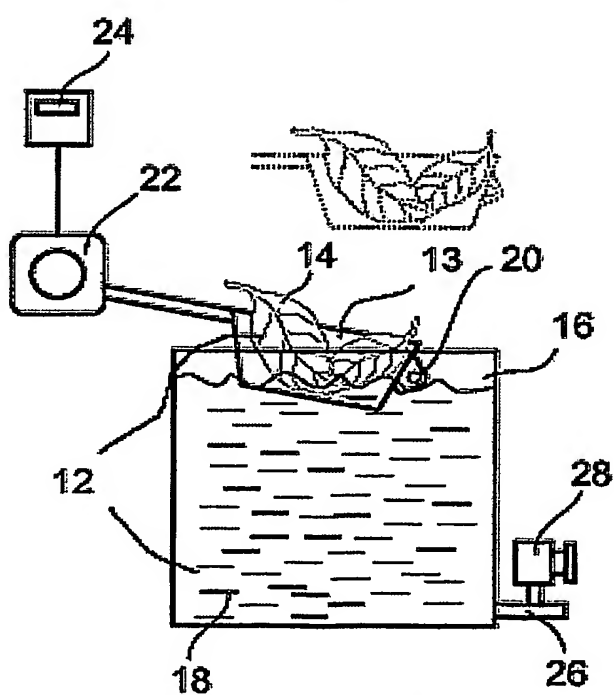


Fig. 4b

10

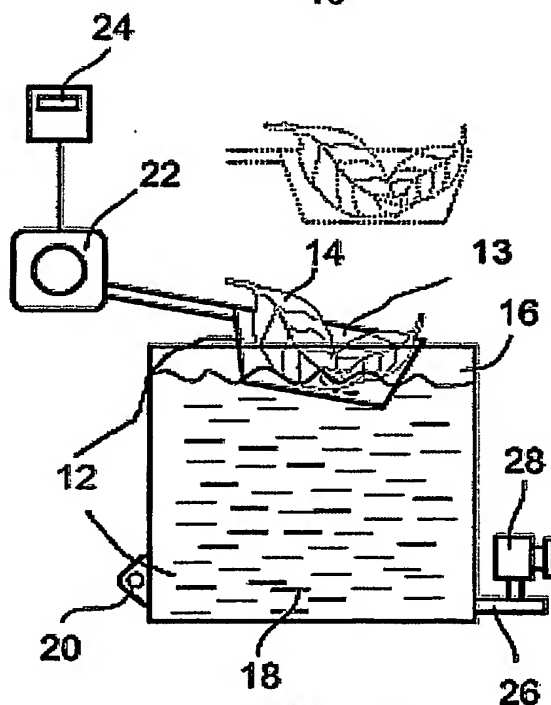


Fig. 4c

12/17

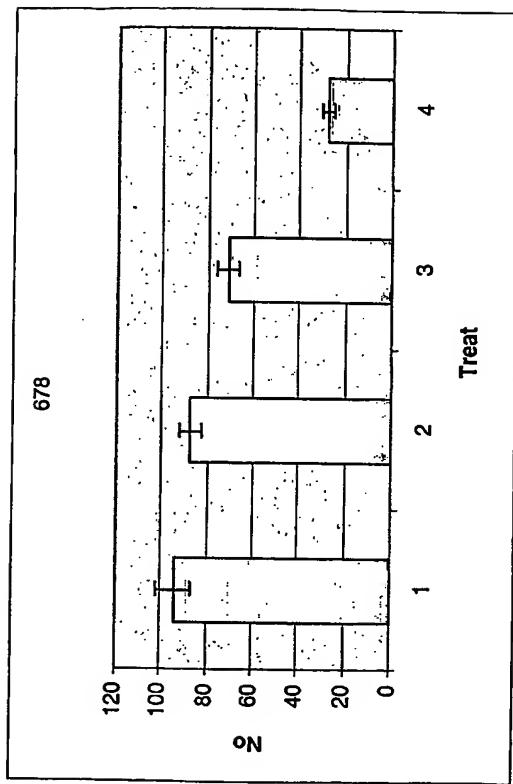


Fig. 5a

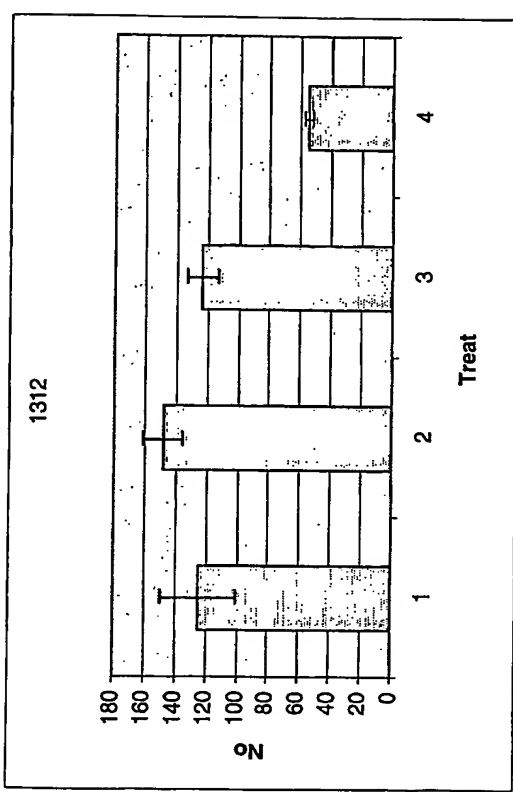


Fig. 5b

13/17

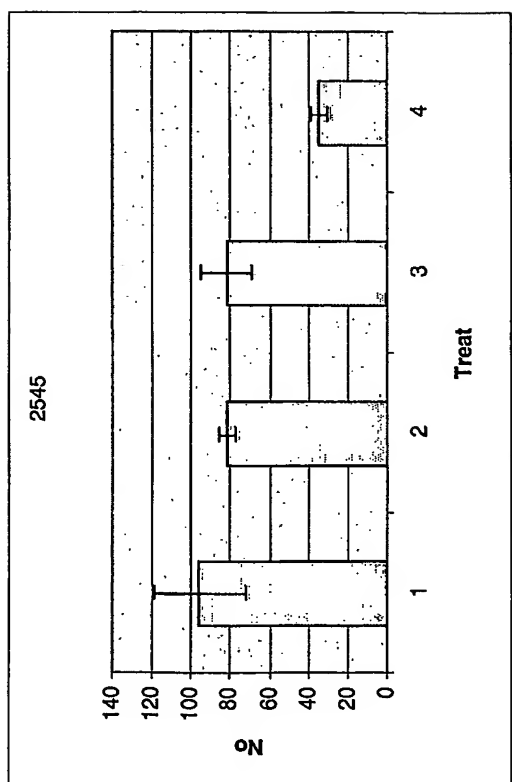
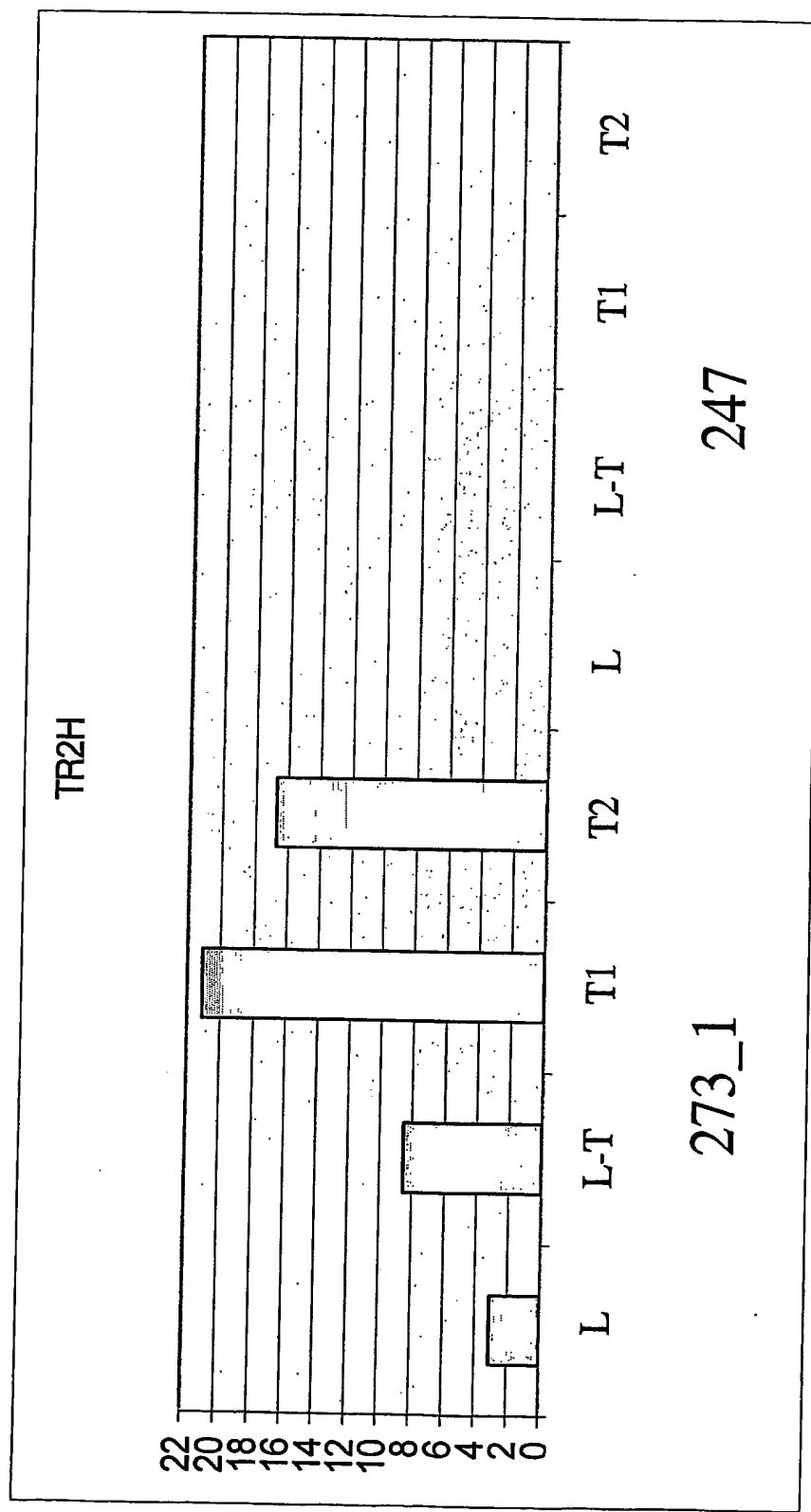


Fig. 5c

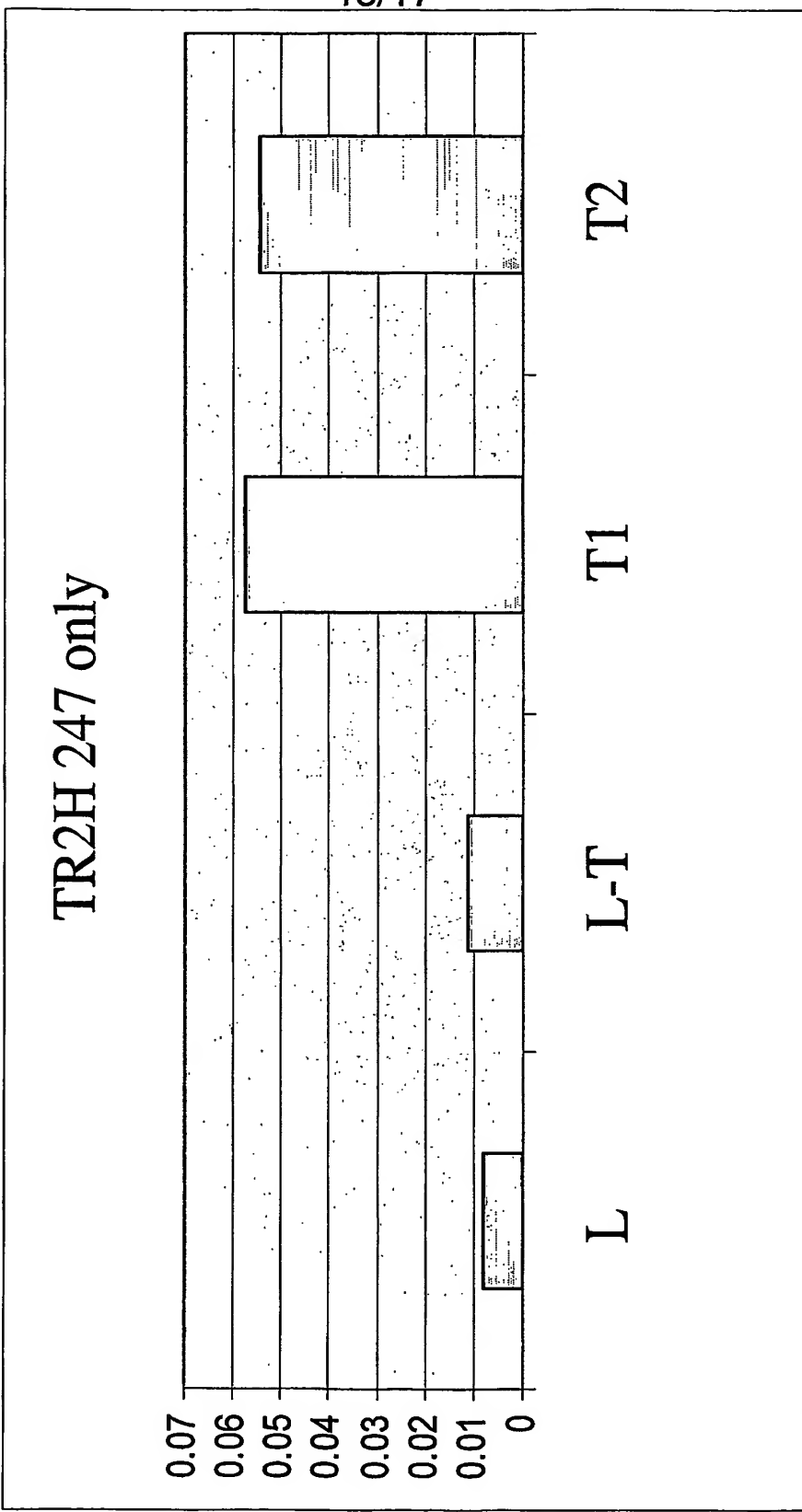
14/17

Fig. 6a



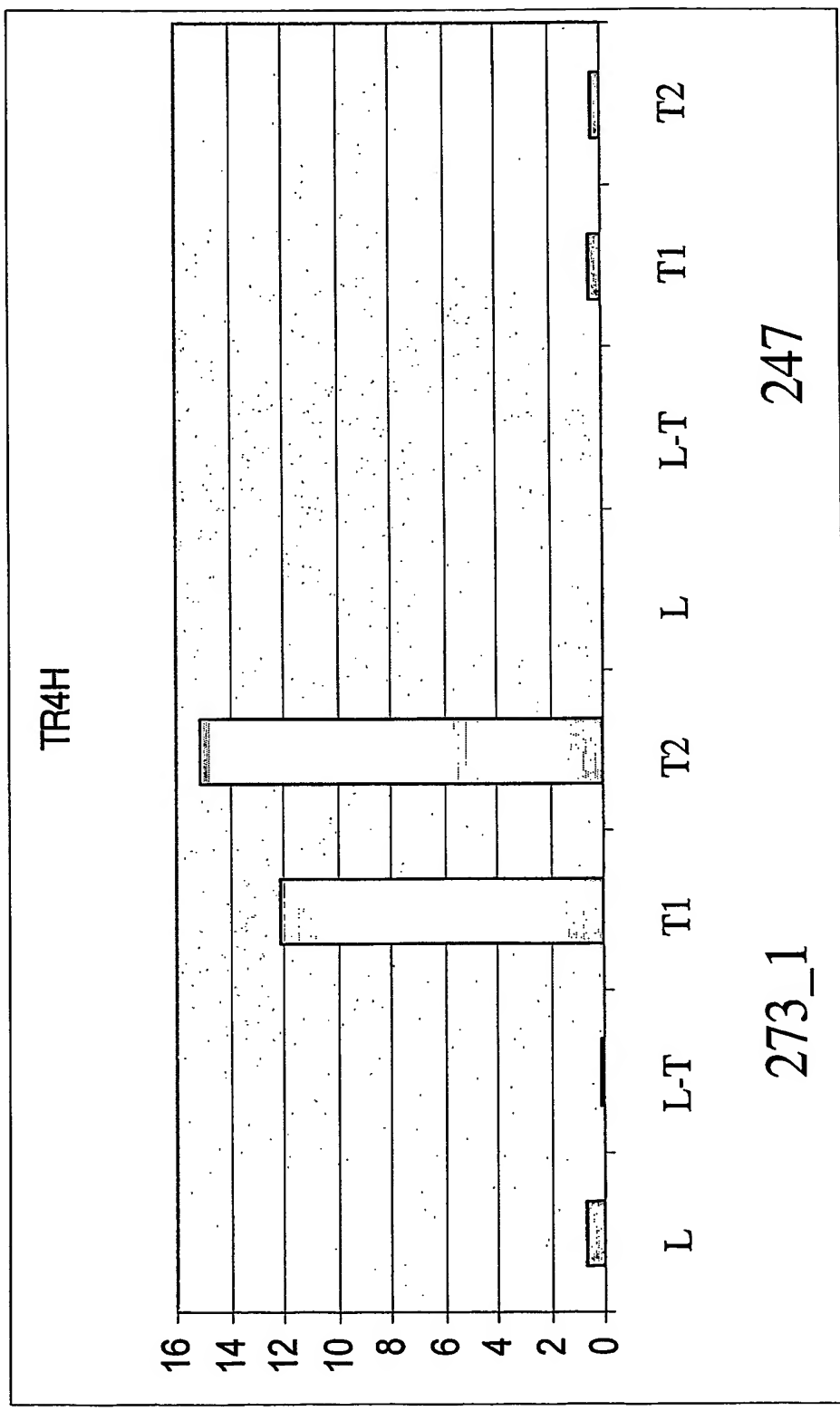
15/17

Fig. 6b



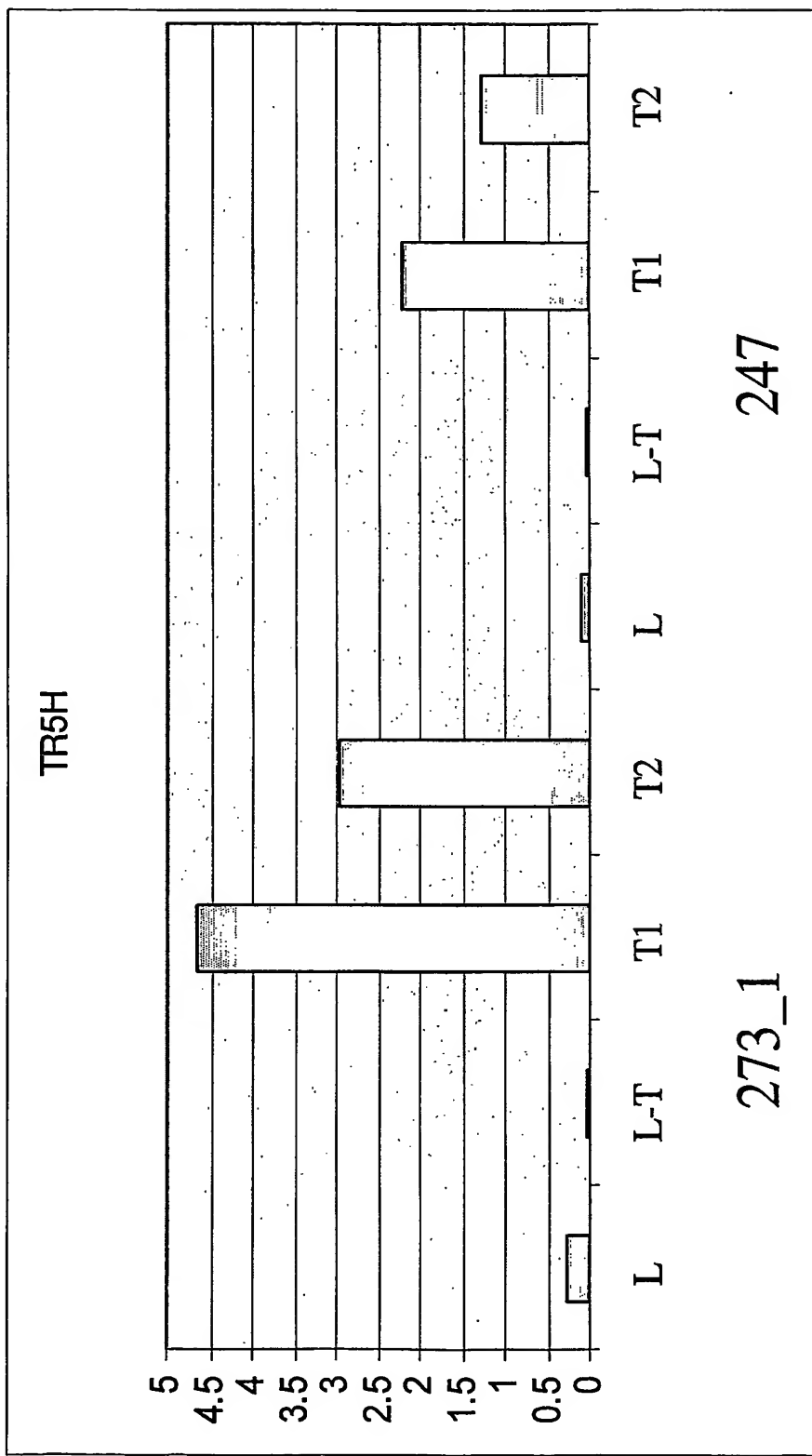
16/17

Fig. 6c



17/17

Fig. 6d



1
SEQUENCE LISTING

<110> EVOGENE LTD.
Ronen, Gil
Rabinovich, Larisa
Meissner, Rafael
Karchi, Hagai

<120> NUCLEOTIDE SEQUENCES FOR REGULATING GENE EXPRESSION IN PLANT
TRICHOMES AND CONSTRUCTS AND METHODS UTILIZING SAME

<130> 27120

<160> 88

<170> PatentIn version 3.2

<210> 1
<211> 26
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 1
atggaagtaa ctttgttgt tagtac

26

<210> 2
<211> 21
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 2
gccagtgtatc accataagga g

21

<210> 3
<211> 376
<212> DNA
<213> Gossypium hirsutum

<400> 3
atggaagtaa ctttgttgt tagtacttca ctctctattt tgtttgct tctacttgtt
aaaccttgttt catcaaaaacg aagaaaaacag aatctaccac caagcccaact acttaaaactt
60
120

ccaatattag gccatctcta tcccttaaa ccacmtctat atcgactct tgctaatctc	180
tcaactaaat atggccctgt ttctcttta caattaggta cccgtttgt tgttagcaatt	240
tcctcaccat ctgctgccga agaatgtttc aaaaaaaatg atatcgaaaa tgctaatgc	300
cctcggacaa tgacggcaaa attcataggc tataactcta ctacagtcat tggttctact	360
tatggtgatc actggc	376

<210> 4
<211> 22
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 4
ttctttgggtt cttcaatgtt gg 22

<210> 5
<211> 22
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 5
tttgaatgt cattgggagg tc 22

<210> 6
<211> 410
<212> DNA
<213> *Gossypium hirsutum*

<400> 6
ttctttgggtt cttcaatgtt ggaaattatc atctccatct ctgattttac aacaaaatc 60
atcaatatcc atgggtgcat tcaaaggta attacttatg ggaattttt aattttgttc 120
ataccttata tacgtacaca tgaaaaaaatt gactattaaat tttgttaggta ttcataaaact 180
tcaaatcccc aattccatcc tgacagtgtc tgctcgtgga ctcacaaaga tttcatgctc 240

3

actcaactta caaacggaaa agctttgtta tgaggataat gataatgatc ttgatgaaga	300
acttatgcct aaacacattt ctttgataat ggatggtaat aggagatggg caaaggataa	360
gggttttagaa gtatatgaag gtcacaaaca tattattcca aaattaaaag	410

<210> 7
<211> 23
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 7
gggtaatatt catttgattt tcc

23

<210> 8
<211> 22
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 8
aacctgcattt acatgtttca ag

22

<210> 9
<211> 431
<212> DNA
<213> *Gossypium hirsutum*

<400> 9
gggtaatatt catttgattt tcccactttt atttatatct tgtttcattt tccccatccac
60
aacaaatggc tactccaacg caatcataaa agcttgggc acccaaacac ctcatccaca
120
accttgcattt tacttcttat cacaatcc caaaattaca ttccttatca taaaaaaatc
180
agattttcta aaagtgtcac tagacttagt gtttagaccgt gcgttacgtc cccaaactgaa
240
cacatattca ctaggtccaa aatgtcgtaa cgagcgcgaa aaaaacgcgt gggctgattt
300
cattgaactc tatgaaaact caatcaacaa aatcaaaagc acagttgatc caaacacaaa
360
atgctcagct actgatgctc aaacatggtt aagtacatcc ttaacaaatc ttgaaacatg
420

taaagcagg t

431

<210> 10
<211> 28
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 10
ttttttttt tgtttgttgt ggggggtgt

28

<210> 11
<211> 22
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 11
ggaagttaa gtatgtggct tg

22

<210> 12
<211> 17
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 12
tttttgtttg ttgtggg

17

<210> 13
<211> 20
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 13
gtgggcttgg tggtagatcc

20

<210> 14
<211> 3284
<212> DNA
<213> *Gossypium hirsutum*

<400> 14	
tttttgttg ttgtgggggt gtcgctcagc ccctactcat ccaagggtt a	60
aaaataaaaa taattgcate aattttaaaa aaaaaaagat ggcaaattga caatcatatt	120
aaagatggta ggtgttaactt cactacatta tttattgtgt ctttaagag ctcaaatcct	180
ttgctttctt aaataaaaat aaaaaaagca agaaaattat actaactcct atttcacagg	240
gcctccattg aagtctttt attggtagt tgataaagt ataaaccgaat gactaagagc	300
ctgtttggat cagttaaaaa gctggtcaaa ctgacttaca agctgattt tgacttattt	360
agctgttga caatactgaa aataacttat ttaagttaa aaaaaaata ttatTTtaag	420
ccaaaagtta aaagtgggg gagaggtgt tttttttt agttataag ttgttttaag	480
ttgaccacat tttatgttt ttgccttaa tttttata caatctccaa attagaacat	540
aaccctaaca tctctttttt ccattttcc ctttcacgt ttgacatagc aacctcagca	600
cttttatcca aacacataac tgctatattt taaaaataag tttcagcact ttcaaaagta	660
ctttttaaa gttgctttta ttaagccat ccaaacgcgc cctaataaat ctctttact	720
ttgtcgata ttagctctat attcaacaa atatagttt tctttattct taacgtattc	780
atgttctttt caatttgtct tatttattac tattatatga ttatagttt ttatacatat	840
gatatgttc gtctagagta agtcatgttt tatctagaat aagtctatTT taaaacaaaat	900
gtaattcaat tagtagaaa atatTTTct ctatataattt taatgtatg tctttttt	960
ggatgccatg tctttatTTt cttttttttt cgatttgaaa ttgtatTTt tttataaat	1020
ttgatggat accgctcaaa cattttgtaa ttttttatt tatacgtatc ttttttata	1080
gacaatttat ttttgatta taaaattta tgtttagtaa taaaatatg ttaattcctc	1140
tgataaaaata aatgtttata tttcatgaag tattcaatat atcagacctc caacatctaa	1200
cacaagattt tcattttata tttgtgtaa agttatattc ttatgttaaa cttacatacg	1260

aaaggattta gat~~ttaaact tagcttatata aattaaaatt ttctaatate aattagggga~~ 1320
taaacgtgtg atgcacgcac gttccgagaa ttagttatta ttattaatat atgaagtctt 1380
attgatcaaa aatcacgcac gttccgagaa ttagttatta ttattaatat atgaagtctt 1440
attgatcaaa aagaaaaaaa ctcacaaaat acgccaacgc atactttcta ttttaatacg 1500
ctttgcata~~g ataaaaatat ttgttaggatt ttgtgttaact actattagtc cattactatg~~ 1560
ac~~tttattgtg aaaagtgaaa acatgattt tacaaaagaa tctcttaata aaatttattg~~ 1620
attattat~~tttcttaggc ggggaaaat aagtagttt ataaatattt ttttaagaat~~ 1680
ttgtgattt taattgtcg aaagttaaa ccttata~~gat agttacatc tcataat~~ 1740
tttaccacta ttgcaatatt ttcata~~tcta aactatgctt ttctatgaat ttcttaatt~~ 1800
cttttaaatt ttcttaaaat cttaatata~~t tttctacata tttgtatta tattataat~~ 1860
ttaaaaat~~at aggggtcat ggcttacgtt gcttttcttg gtcatactt gattggttct~~ 1920
agaagat~~gta gatgtatcta tcttggcata caaggctaca aagcagccag agagtcctcg~~ 1980
gaattttat ttttttact tttcattttt gaaaaaagta agaaagtaca tatattttt 2040
ttcttattac acttttgac atatttgat tgcattaagg tcaagtaaaa aagtgataac 2100
taatccaaa gagagagtag taatcataca gaaaaaattt attacctacg ggatataatt 2160
attatcagtt gtatgaggct tatttagctg ccacatatta aaaagactca ccttca~~ctt~~ 2220
tcatttcatt tccttatctc ttttattta accttttct tctgtacttt tactctctc 2280
ccaaactctt cttgttttt tctat~~ttgtt~~ attaacattt aatataattt tattttttc 2340
aatccgacat ttgcattaaa attagaat~~at tttaaattta aaattgtgta aggctttatt~~ 2400
caaagaaata tataatctat caaaaaagaa tttcatattc aaaatttgaa ctcgagactt 2460
ctaattaagt aagaataaa tatcatcctg tactccatca tatatttgag gaaaaaccaa 2520
aataggtgtg tagaaatatt taaaatta~~t tttttggat gagtttaag gaattgaaga~~ 2580
aagtgc~~aaca aaaaaata ataattgtga aattaattt ttgttttgc atttatttc~~ 2640
taatttgatt ttttgaata atatcaaag tgcactttat atatataaaa ctcattaaac 2700
aattaaattt gaattttaa ctattcatca aacaattgat ggggttcttg cttaactaga 2760

7

ggttttaagt ttcaaatttt aaatacaaaa aattcttggt gacaacatat aatcgaaat 2820
taacacaaaat attgaatata aaataaaaata taagaagagt taataagtag gggaaaaat 2880
gaagaacagt tgggaggaaa aagacggtaa aaaaaagggt taaaaatgaa agaaaaagga 2940
aatgtaatga aatgaagtga aatatggatc ccattaacac gttgcagcca aacaaggcct 3000
tatacaaccg acaataatta tatctcgctt aaaataaaat tttttgtatc acgcgtata 3060
aatttgaacc aatattttct tgagtggacc cataagttga aaagtctagg ctggttcaac 3120
agccccatca tctatactat tataatataaa ccaattcagt gcaacaagtt gagatatgga 3180
agtaactttg ttgtatagta cttcactctc tattttgttt gtgcttctac ttgttaaact 3240
tgtttcatca aaacgaagaa aacagaatct accaccaagc ccac 3284

<210> 15
<211> 21
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 15
gtttagtcca cgagcagaca c 21

<210> 16
<211> 20
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 16
cgagcagaca ctgtcagagg 20

<210> 17
<211> 1337
<212> DNA
<213> Gossypium hirsutum

<400> 17
ttttgtttg ttgtgggggt gttaaatggt gggttgggt gaaattggaa atattacaat 60

gggtttgaat agaaattggg ttgggttaga cccgcccaaa tttactttaa actcaaatga	120
gctaaaaata ggttggcct tgacccgccc aatttgcatt gattaatctt agttatctttaa	180
catattgata tttaactttt ataatcacat tttgaaggtc cgttcaagaa tttttgtta	240
aaaaaaagttaa caaatggata gataaatcat aaaaaaggca acaaatcgat aataatttat	300
attgtaaata taggaacata tcttaatact aagttctaaa acgggttcaa attggagatt	360
gaatttaggct taattgagaa ttcttttcaa ataggtaag cttgaatggg tcgagattga	420
acccaattca aattatcttg agcccaaccc ttaaaattct gggegaattt ggcattttac	480
catgtttggg ttcattttta acgcccctag cgtagtcgaa agaagtcaat ccatgagggtt	540
tgtaaaaaca atgccaataa ttaccctac cattgagctt gtttgtcatg tggtagca	600
aaatggtaga ttatcgaaaa aatatcttaa ttatgttca tagttataat ttgttaatta	660
caatttagtag ctacatgtta tatggaggag agtggtgagc gagattggg gaggaaagag	720
agaagtgagt gagacaaggt agagagtggg agagaggcga actgcataatg catatttgc	780
aaaataattt tatatatgtt actggatatac atacgtattt gtatatctgg tgagtggg	840
gagaaaaagag agaagcgagc gagattggaa gaggaaagag agagccgagc gagagaggac	900
aataattttt gtaattcgca tctcatttgcataattt ttgttgcataa tgccgttcaaa	960
tataattttt taaccataag cataaacaac ccttatataga actattgttca aatatagaac	1020
tattgtatcta ttgatcaaaa gagtcataacc ataattctat tttaaacacca cctcccttgt	1080
ttcacttcac aataaaataa atttgagtaa taaagcatga gttttttgtt tttcaatgt	1140
tggaaattat catctccatc tctgattttta caacaaaata catcaatatac catgggtgca	1200
ttcaaagggtt aattacttat gggattttttaatttttgtt cataccttat atacgtacac	1260
atgaaaaaaat tgactattaa tttttaggtt attcataaacc ttcaaatccc aaattcgctt	1320
ctgacagtgt ctgtcg	1337

<210> 18

<211> 22

<212> DNA

<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 18
attcacaagg ttgtggatga gg

22

<210> 19
<211> 20
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 19
gatgaggtgt ttgggtgcac

20

<210> 20
<211> 1906
<212> DNA
<213> Gossypium hirsutum

<400> 20
tttttgttg ttgtgggggt gtcattgagt ctttcaagg tgtgaatctt ttaacgaaaa 60
gacctgctct gataccaatt gaagaaacct tacccagaa cacgaaccag gttcggtgtaa 120
gttgctttta agtaaagaca gagtaaagac acaaacactt attgaattaa aaaccttcct 180
cgctcaagga aggaaaaacc tcgttttatt aattcaacta taagatttg tgattacaac 240
tcaataatca aaaagtctta tctctactac tccctcgatt gactccaatc gatctctcca 300
aaagggtcaaa cccacctttt gttacaattc tcacagaaaac tcaaccctac aaagagccaa 360
acccactcct tgtacaactc tcacagaaaac acaaccctac aagaagtcaa acccactcct 420
tgtacaactc tcacagaaaac tcaaccctac aagaagccaa acccactcct tgtacaataa 480
ctcgtaactt acaatcaaga acgaaacaag aagatagttt tacacgttga aaaccttctc 540
actcaagaat gttttaaacg tagtaatcct atcaaccttg aagacttcaa tttgataaat 600
aattctccct tggctctgc gtgaagtctg cgtttcttc ctctgoctcg tgctttttt 660
atagagtttg tttgccttg tgcaatcctt tttgataagg taaggaagtt atgtttaaac 720

10

aagaattccc	ttttaaagta	caatccttat	tatataacaac	ttcgttcctt	aataatataat	780
ttaagggttt	ccttatttgt	atcaacttat	accttaataa	tattatttt	ggcttgaca	840
aataactcta	ttttcttgat	tactggctg	accactta	ctcgatcttgc	gactcgagct	900
tggcttcctt	tgctgcgtac	atttgtact	gattattgc	gcttcgttc	tatcatcaaa	960
acatgaatta	tcgattcaat	catattctat	cagctactat	ttagttggaa	tgtttgagaa	1020
cacacaaaag	ttttcaaaa	cttgaactga	aatgtctaat	aaaaacactc	tatctatcat	1080
attttagat	ctcaattgaa	ataacatatt	atgattcgat	tctctaaaaa	taaaaatttc	1140
gtagcttaa	gagattatta	atatattaag	tgataattta	atgttagtta	attagttaaa	1200
atctaacgtg	tggtaggtaa	tacatataagg	aatacgccct	ctctagcttc	ctgtttcca	1260
cttttaaag	ttgggttcctt	gttcatcag	tttaattcc	ttatcaagtc	atcaaacaca	1320
cataattacc	cgcagaattt	taattttttt	ttaatttatta	cattatgat	tagattttt	1380
tcttccaaaa	cctaagaaat	agccacacac	gtatggttct	cactattcat	gccttaagga	1440
aaaaaaataa	aaaagaggat	ggtgcateccc	catcaacttag	tttttgacat	tccgttgac	1500
ctcttatatt	cctatatcta	tataaagaac	ccaaaagaca	ccaaatacaa	tcacagtctc	1560
tctcaaaaaa	aaaaaacata	ttacaaactc	cttacgatgg	gtaatattca	tttgattttc	1620
ccacttttat	ttatatcttgc	tttcattttc	ccatccacaa	caaatggcta	ctccaacgca	1680
atcataaaag	cttggtgac	ccaaacaccc	catccacaac	cttgtgaata	cttcttatca	1740
ccaaatcccc	aaattacatc	tcctatcata	aaaaaattcag	attttctaaa	agtgtcacta	1800
gacttagtgt	tagaccgtgc	gttacgtgcc	caactgaaca	catattcact	aggcggaaaa	1860
tgtcgtaacg	agcgcgaaaa	aaacgcattgg	gctgattgca	ttgaac		1906

<210> 21
<211> 31
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 21

11

aatttaagct tgggtcgctc agcccccact c

31

<210> 22

<211> 35

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 22

aaatttgcga catctcaact tgttgcactg aattg

35

<210> 23

<211> 3157

<212> DNA

<213> Gossypium hirsutum

<400> 23

gtgtcgctca gcccctactc atccaagggt tattcaaaaa ataaaataaa aataattgca

60

tcaattttaa aaaaaaaaaag atggcaaatt gacaatcata taaaagatgg taggtgtaac

120

ttcactacat tatttattgt gtcctttaag agctcaaattc ctttgctttc taaaataaaa

180

ataaaaaaaaaag caagaaaatt atactaactc ctatccaca ggccctccat tgaagtcttt

240

tgatttggtt gttgataaaag ttataaccga atgactaaga gcctgtttgg atcagtttaa

300

aagctggtca aactgactta caagctgatt tttgacttat ttagctgttt gacaatactg

360

aaaataactt attttaagtt aaaaaaaaaaa tattatTTTA agccaaaagt taaaagttgg

420

gggagaggtg cttttttttt ttagcttata agttgtttta agttgaccac atttttatgt

480

ttttgccctt aatatTTTA tacaatctcc aaatttggAAC ataaCCCTAA catcttttc

540

ttccattttt ccctttcac gtttgcata gcaacttcag cacttttac caaacacata

600

actgctatat tttaaaaata agtttcagca ctttccaaag tacttttta aagttgttt

660

tattaaggccc atccaaacgc gccctaataa atcttttaa ctttgtgttta tattagctct

720

atatttcaac aaatatagtt tatctttatt cttAACGTAT tcattttttt ttcaattttgt

780

cttattttatt actattatat gattatagtt ttttatacat atgatatgtt tcgtctagag

840

taagtcatgt ttatctaga ataagtctat tttaaacaaa atgtaattca attagttga

900

aaatattttt ctctatataat ttaatgtaa tgtcttttt ttggatgcca tgtctttatt	960
tacttccttt tcgcatttga aattgtatt tttttataa atttgatggt ataccgctca	1020
aacattttgt aatattttta ttatcacgt a cttttttta tagacaattt attttttgat	1080
tattaaaatt tatgttttagt aattaaaata tgtaattcc tctgataaaa taaatgttta	1140
tatccatga agtattcaat atatcagacc tccaacatct aacacaagat ttcatgtta	1200
tattttgtgt aaagtatat tcttatgtta aacttacata cgaaaggatt tagatttaaa	1260
ccttagctata taaattaaaa ttttctaata tcaatttaggg gataaacgtg tgatgcacgc	1320
acgttccgag aattagttat tattattaat atatgaagtc ttattgatca aaaatcacgc	1380
acgttccgag aattagttat tattattaat atatgaagtc ttattgatca aaaagaaaaaa	1440
aactcacaaa atacgccaac gcatacttcc tattttataa cgctttgcat agataaaaat	1500
atttgttagga ttttgtgtta ctactattag tccattacta tgacctattg tgaaaagtga	1560
aaacatgatt tttcacaaaag aatctcttaa taaaatttat tgattattat ttctttctag	1620
gcgggggaaa ataagtagtt tgataaatat ttttttaaga atttgtgatt tttaattgtt	1680
cgaaagttaa aaccttatacg ttagcttaca tctcatatata attttaccac tattgcaata	1740
ttttcatatc taaactatgc ttttctatga atttcttaa ttctttaaa ttttcttaaa	1800
atcttaatata ttttctaca tattttgtat tatattataa atttaaaaat atagggggtc	1860
atggcttacg ttgcctttct tggctcatc ac ttgattggtt ctagaagatg tagatgtatc	1920
tatcttggca tacaaggcta caaagcagcc agagagtcct cgaaattttt atttttttta	1980
cttttcattt ttgaaaaaaag taagaaaagta catatattt ttttcttatt acacttttg	2040
acatatttgtt attgcattaa ggtcaagtaa aaaagtgata actaaatcca aagagagagt	2100
agtaatcata cagaaaaaaat ttattaccta cgggatataa ttattatcag ttgtatgagg	2160
cttatttagc tgccacatata taaaagact caccttcac ttcatatca ttcttattatc	2220
tcttttattt taacctttt cttctgtact tttactctcttccactctt ttcttgcattt	2280
tttcttatttgc ttatataacat ttaatataat tttatatttt tcaatccgac atttgcattt	2340
aaatttagaaat attttaaattt taaaattgtg taaggctta ttcaaagaaa tatataatct	2400

atcaaaaaag aatttcatat tcaaaatttg aactcgagac ttctaattaa gtaagaata	2460
aatatcatcc tgtactccat catatatttg agaaaaacc aaaataggtg tgtagaata	2520
tttaaaatta attttttgg atgagttta aggaattgaa gaaagtcaa caacaaaaaa	2580
taataattgt gaaattaatt ttttgtttt gcatttattt tctaatttga ttttttgaa	2640
taatatcaaa agtgcacttt atatatataa aactcattaa acaattaaat ttgaatttt	2700
aactattcat caaacaattg atggggttct tgcttaacta gaggtttaa gtttcaaatt	2760
ttaaatacaa aaaattcttg ttgacaacat ataatcgaat tttAACACAA atattgaata	2820
taaaataaaa tataagaaga gttaataagt agggaaaaaa atgaagaaca gttgggagga	2880
aaaagacggt aaaaaaagg gttaaaaatg aaagaaaaag gaaatgtaat gaaatgaagt	2940
gaaatatgga tcccattaac acgttgcagc caaacaaggc cttatacaac cgacaataat	3000
tatatctcgc taaaataaa atttttgtt tcacgcgtaa taaatttcaa ccaatattt	3060
cttgagtgga cccataagtt gaaaagtcta ggctggttca acagccccat catctatact	3120
attatatata aaccaattca gtgcaacaag ttgagat	3157

<210> 24
<211> 30
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 24
cctagtcgac ggtgttaat ggtgggttgg 30

<210> 25
<211> 27
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 25
ttggatccga gcagacactg tcagagg . 27

<210> 26
 <211> 1320
 <212> DNA
 <213> *Gossypium hirsutum*

<400> 26	
ggtgttaaat ggtgggttgg gttgaaatttg gaaatattac aatgggtttg aatagaatt	60
gggttgggtt agacccgccc aaatttactt tgaactcaaa tgagctaaaa atagggttgg	120
ccttgacccg cccaatttga tccgattaat cttagttatt taacatattt atatthaact	180
tttataatca cattttgaag ttccgttcaa gaattttttg ttaaaaaaaaag taacaaatgg	240
atagataaaat cataaaaaaag gcaacaaatc gataataatt tatattgtaa atataggaac	300
atatcttaat actaagttct aaaacgggtt gaaattggag attgaatttag gcttaattga	360
gaattctttt caaataggtt aagcttgaat gggtcgagat tgaacccat tcaaatttac	420
ttgagcccaa cccttaaaaat tctggcgaa ttgggcatgt taccatgttt gggttcattt	480
ttaacgcccc tagcgtagtc gaaagaagtgc aatccatgag gtttgtaaaa caaatgcgaa	540
taatttaccc taccatttgcg cttgttagtc atatggtta gcaaaatggt agattatcga	600
aaaaatatct taattatgct tcatagttat aatttgttaa ttacaatttag tagctacatg	660
ttatatggag gagagtggtg agcgagattt ggagaggaaa gagagaagtgc agttagacaa	720
ggtagagagt gggagagagg cgaactgcattt atgcattttt gtcaaaataa ttgtatata	780
gtaactggta tacatacgta ttcttatatc tggtgagtga ggagagaaaa gagagaagcg	840
agcgagattt gaaaggaaaa gagagagccg agcgagagag gacaataatt tatgttaattc	900
gcatctcatt tgtataatta attttggcg aaatgcggtt caatataatt ttttaaccat	960
aagcataaac aaccctatat agaactatttgc atcaatatacg aactatttgcat ctattgtca	1020
aaagagtcattt accataatttgc tattttaaaca ccacccctt tgtttgcattt cacaataaaa	1080
taaatttgag taataaagca tgagttttt gggttccaa tggtggaaat tatcatctcc	1140
atctctgatt ttacaacaaa atacatcaat atccatgggt gcattcaag gttaattact	1200
tatggaaattt atttaattttt gttcatacctt tataacgtt cacatggaaaa aatttgactat	1260

15

taattttgta ggtattcata aacttcaaat cccaaattcg cctctgacag tgtctgtcg 1320

<210> 27

<211> 32

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 27

tttccaagct tgacctgctc tgataccaat tg

32

<210> 28

<211> 29

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 28

ccggatccctc gtaaggagtt tgcataatgt

29

<210> 29

<211> 1537

<212> DNA

<213> Gossypium hirsutum

<400> 29

gacctgctct gataccaatt gaagaaacct taccccaagaa cacgaaccag gttcggtgaa

60

gttgctttta agtaaaagaca gagtaaagac acaaacactt attgaattaa aaaccttcct

120

cgctcaagga aggaaaaacc tcgttttatt aattcaacta taagattttg tgattacaac

180

tcaataatca aaaagtctta tctctactac tccctcgatt gactccaatc gatctctcca

240

aaaggtaaaa cccacccccc gttacaattc tcacagaaac tcaaccctac aaagagccaa

300

acccactcct tgcataactc tcacagaaac acaaccctac aagaagtcaa acccactcct

360

tgcataactc tcacagaaac tcaaccctac aagaagccaa acccactcct tgcataataa

420

ctcgtaactt acaatcaaga acgaaacaag aagatagttt tacacgttga aaaccttcctc

480

actcaagaat gttttaaacg tagtaatcct atcaacccctt aagacttcaa tttgataaaat

540

aattctccct	tgttctctgc	gtgaagtcgt	cgttttcttc	ctctgcctcg	tgcctttttt	600
atagagtttg	ttttgccttg	tgcataatcctt	tttgataagg	taaggaagtt	atgtttaaac	660
aagaattccc	ttttaaagta	caatccttat	tatataacaac	ttccttcctt	aataatatat	720
ttaaggaaaa	ccttattttgt	atcaacttat	acctttaata	tattttttt	ggctttgaca	780
aataactcta	ttttcttgat	tacttggctg	accacattt	ctcgatctt	gactcgagct	840
tggcttcctt	tgcgcgtac	atggctact	gattatttgc	gcttccttgc	tatcatcaaa	900
acatgaatta	tcgattcaat	catattctat	cagctactat	ttagttggaa	tgtttgagaa	960
cacacaaaag	tttttcaaaa	cttgaactga	aatgtctaat	aaaaacactc	tatctatcat	1020
attttttagat	ctcaatttgc	ataacatatt	atgatcgat	tctctaaaaa	taaaaatttc	1080
gtagctttaa	gagattatta	atataatgg	tgataatttta	atgttagtta	attagttaaa	1140
atctaacgtg	tggtaggtaa	tacatataagg	aatacgcct	ctctagctt	ctgttttcca	1200
ctttttaaag	ttgggttcctt	gtttcatcag	tttaattttcc	ttatcaagtc	atcaaacaca	1260
cataattacc	cgcagaattt	taattttttt	ttaatttatta	cattatgtat	tagattttt	1320
tcttccaaaa	cctaagaaat	agccacacac	gtatggttct	cactattcat	gccttaagga	1380
aaaaaaaaataa	aaaagaggat	ggtgcattcc	catcaattag	tttttgacat	tccgttgtac	1440
ctcttatatt	cctatatcta	tataaagaac	ccaaaagaca	ccaaatacaa	tcacagtctc	1500
tctcaaaaaaa	aaaaaacata	ttacaaactc	cttacga			1537

<210> 30

<211> 30

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 30

aaatcttagac taccatcgct agtaatcggt 30

<210> 31

<211> 24

<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 31
gttgaagaac tgcatccgg gagg

24

<210> 32
<211> 1834
<212> DNA
<213> Nicotiana tabacum

<400> 32
tctagactac catcgctagt aatcggtggca ataactaccc taaactatagc atttattgtct 60
accaaataaaa atttggcagc taatcataat tttttgtcat gaatcaatag ttatttgtac 120
aatagtttac tcttagccac aataaattat ttaaaataaaa atattatagc taaataaata 180
tttttgcttt aagttctaaa agcttgtggc aatagttaaa tgatatagtc acagatttat 240
tggtataatt gaattatgtt gctaatttct tagtttttg ccacgagttttaaaaattacca 300
atagctatag taactttta atcacaataa aatatttgaa agaaaatattt gtagctaaat 360
gaatattttt tccttcaagt tattaaaagt tgtggcaata taggttaaat tagccacatg 420
tttcttgctt taatagaatt ttgttagctaa tcattaactt ttaccacgag ttgaacttaa 480
tataacaaca ataaccctttt aaccataata aagcgattta aatcaaataat tactaaataa 540
ataactttgc tttcaagttt ctataaaatc atggcaatag tcattacgat aaaatgatat 600
aaccacgaat atattgcaac gataaattct gtaactaatac attagttttt gcgacgaggt 660
aaattttccg tcacagtagc aatcttctag gcacattaaa aatttgaacaaaatttgt 720
agtcaaataaa atatttatct tcttatttttta agaaaataaaa aatagtttaga taatagttac 780
tactatttgtt catgaaaata tcaatagata caaattttaaa gtgactataa attacgagt 840
ttactataact ttagtcgtac agtttgcattt aatagtattt taaccacaat tagttatatg 900
tacaaaataaa cataagtgaa taactttttt tcaatgagaa aataagagtt gctcaaacaa 960
tatcaagtttta caaaaattta attttaactg taaaagttat atttttccaa aataacataa 1020

18

actatagtaa ttatataatg tttgaagtat taataaaatt taaatatgca aaagttattt 1080
ttaataaacc atttgtatgc ctaacttgta gcctctaaac tattttatgg cttttatgg 1140
tc当地actcat attttattttt attgcacccctt gtttagtttg gacgttaattt atatataat 1200
ggtgtaaaat ttaaaaatata ttaacatgg tggagaattt atgtatgcct gggttcttaac 1260
tattttttt tatataactg gttagagtaa ttctttat ttcagtatgg atttttaaat 1320
aagtccctcat aaattgaaga cttaaaaatgt ttttgggtca ttccctttttt tatttaagaa 1380
attgaagaat tccgctaaat ttcatatggc cgctgttattt taactgtttt tttcccttgg 1440
taatataattt ggtaagaagt tttaaaataa aggagttat gattttcttag gttcatggct 1500
tgccttagttt ctacgagtaa gogccatcac gactcccgag gataaggaaa tccgggtcgt 1560
agcattcaact cacaaaaattt actaaaaaca aagtttaccc ttctcccaaa agtaaatttc 1620
atatttggct ccacataatg tggtcaatga gtcaagtggaa gtacttttca tgacaaaaaaa 1680
aagttgctga aaaatgcata tctcatatgg tttttttaga gaaatccccat ttcttgcccta 1740
aacgaaagcc tataaaagag catatattgc aacaacagtt tgcaaaaaact atcaagtcaa 1800
ataatcccccc cttaattcc ctcccaaacc cggg 1834

<210> 33

<211> 32

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 33

aaatcttagat aagttgataa agctaatttc tc

32

<210> 34

<211> 23

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 34

19

23

tttccccggga cctggaggca atc

<210> 35

<211> 1969

<212> DNA

<213> Nicotiana tabacum

<400> 35

tctagataag ttgataaaagc taatttctca ttttagctac catcgctagt aatcggtggca 60

ataactaccc taactatagc atttattgtc accaaataaa atttggcago taatcataat 120

tttttgtcat gaatcaatag ttattgtgc aatagttatc tcttagccac aataaattat 180

ttaaaataaa atattatagc taaataaaata tttttgtttt aagttctaaa agcttggc 240

aatagttaaa tgatatagtc acagatttat tggtataatt gaattatgtt gctaatttct 300

tagtttttg ccacgagtta aaaattacca atagctatag taactttta atcacaataa 360

aatatttgaa agaaaatatt gtagctaaat gaatattttt tccttcagg tattaaaagt 420

tgtggcaata taggttaaat tagccacatg tttcttgctt taatagaatt ttgttagctaa 480

tcattaacctt ttaccacgag ttgaacttaa tataacaaca ataacotttt aaccataata 540

aagcgattta aatcaaataat tactaaataa ataacttgc tttcaagttt ctataaaatc 600

atggcaatag tcattacgt aaaatgatata accacgaat atattgcaac gataaattct 660

gtaactaatac attagttttt gegacgaggt aaattttccg tcacagtagc aatcttctag 720

gcacattaaa aatttggaaac aaaattttgt agtcaaataa atatttatct tcttatttt 780

agaaaataaa aatagttaga taatagttac tactattgt catgaaaata tcaatagata 840

caaattttaaa gtgactataa atttacgagt ttactatact ttagtcgtac agtttgcatt 900

aatagtattt taaccacaaat tagttatag tacaaaataa cataagtcaa taacttttt 960

tcaatgagaa aataagagtt gctcaaacaa tatcaaggta caaaaattta attttaactg 1020

taaaaagttat atttttccaa aataacataa actatagtaa ttatataatag tttgaagtat 1080

taataaaatt taaatatgca aaagtttaatt ttaataaaacc atttgtatgc ctaacttgta 1140

gcgtctaaac tattttatgtt gctttatgtt tcaaaactcat attttattttt attgcacett 1200

gttagttttg gacgttaattt atatataattt ggtgtaaaat ttaaaatata ttaacatttg 1260

20

tggagaat	ttt atgtatgcct ggttcttaac tattttttt tatataactg gtttagagtaa	1320
tttcttat	ttcagtat ttttttaat aagtcctcat aaattgaaga cttaaaagt	1380
ttttgtgtca	ttccctttt tatttaagaa attgaagaat tccgctaaat ttcatattc	1440
cgcgttatt	taactgttta ttcccttgt taatataatt ggtaagaagt tttaaaataa	1500
aggagttat	gatttcttag gtcatggct tgccatgctt ctacgagtaa gcgccatcac	1560
gactcccag	gataaggaaa tccgggtcgt agcattcaact cacaaaaatt actaaaaaca	1620
aagtttaccc	ttctcccaa agtaaatttc atatttgct ccacataatg tgcataatga	1680
gtcaagtgaa	gtactttca tgacaaaaaa aagttgctga aaaatgcata tctcatattt	1740
tttttttaga	gaaatccat ttcttgctta aacgaaagcc tataaaagag catatattgc	1800
aacaacagtt	tgcagaaact atcaagtcaa ataatcccc cttaattcc ctccaaaaat	1860
gcagttttc	aactttttt ccctttcct ttttgtgtca ttctttttt tatttaagaa	1920
atggagaat	tccaatagcc aaacaaaaag attgcctcca ggtcccggg	1969

<210> 36

<211> 32

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 36

tataagcttt aagtttaat cctattgttag tg

32

<210> 37

<211> 25

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 37

cggatccatt aatcacaaga aaaac

25

<210> 38
<211> 625
<212> DNA
<213> *Gossypium hirsutum*

<400> 38

aagcttaag tttaaatcct attgttagtgt tatttataaa aaaaatgaga aaagataaaa	60
atacctttat attaatattt gttatattgt aaaataagga tatttttaac aaattttcaa	120
ttgaatagat gtttgggtga atcctaatac caattaaagt atatatacac aaacaattat	180
aaatcaaatt accttaataa aaatggtatac attcaattca atgacaataa atgcatttat	240
aaatacatca aatgtaaatc tcatgtttat aagaaaacac gtagaaaaaa gttaaaccac	300
tatttgagtc ctagctgtgg aggcatgatt gagtgaaatc aaatggacgc tggttttat	360
tgtattgaaa gaaaccaata atcacgtagg ttggcagttg aacataattt aatggctca	420
acttttaatg tggtgtaat gtttggatcg gataatctca acttacctaa tagctaggaa	480
agtaaaaattc aaacatcacc cgctactact tttggctata aaaaccctcc taccctcaag	540
ccctaaccac gacaatcacc aatagtacta ctactccaag caagtatttt ctttacacgt	600
ttgtttttct tgtgataatg gatcc	625

<210> 39
<211> 627
<212> DNA
<213> *Gossypium hirsutum*

<400> 39

aagcttaag tttaaatcct attgttagtgt tatttataaa aaaaatgaga aaagataaaa	60
atacctttat attaatattt gttatattttt aaaataagga tatttttaac aaattttcaa	120
ttgaatagat gtttgggtga atcctaatac caattaaagt atatatacac caaacaatta	180
taaatcaaatt tacttttaat aaaatgctat cattcaattc aatgacaata aatgcattta	240
taaatacatc aatgttaat ctcatgtttt taagaaaaca cgtagaaaaa aagttaaacc	300
aatatttgag tcctagctgt ggaggcatga ttgagtgaaa tcaaattggac gctggttta	360
attctattga aagaaaccaa taatcacgta gggtggcagt tgaacataat tgaatggct	420
caacttttaa tgtggtgtaa atgtttggat cggtataatct caacttacct aatagctagg	480

22

aaagtaaaaat tcaaacatca cccgctacta cttttggcta taaaaaccct cctaccctca 540
agccctaacc acgacaatca ccaatagtac tactactcca agcaagtatt ttccttacac 600
gtttgtttt cttgtgataa tggatcc 627

<210> 40
<211> 25
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 40
aaaatttggg atctagaagg tgagg 25

<210> 41
<211> 29
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 41
ctggatccta ttgcttagctt tggatgaag 29

<210> 42
<211> 622
<212> DNA
<213> Lycopersicon pennellii

<400> 42
tctagaaggt gaggaacttt ttttaacaat atataagtaa gcattggta taatttcaca 60

acaacattac ggtaaaacct ctataaatta atacccgata aattaataat ccctctaaaa 120

taatattttt ctaggatttt cgatttaggc aatgaaaaaaaa atcaccattt tcaataaaaat 180

aatgagataa tatatttca gaagaccct atataaatac atgggtccta ttaatatcat 240

aaattgatta ttattcaaaa gcataaaatat atctaagata attttagtaaa aaaaatgattc 300

tattctgttt ttttttgtt aaaatttaaa tgttagttgaa gttcatttct aacatttcat 360

23
attgttcca agagctccaa ttttgtctt tcgaactca ccatagaaga gttccagatg 420
cgataagtgt ttcccttacgc gtaattgggtt ccaaaggat agtatcatat tcaacttcat 480
catcgacatt gcttttccg atggatcca taaaattctc taagcttatt tgaaatggag 540
taatatttta tttggccccca acacattata taaggcaatg tatagcccta tgaatcttca 600
tccaaagcta gcaataggat cc 622

<210> 43
<211> 30
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 43
atggaaaago ttatggacag actaaaaacac 30

<210> 44
<211> 30
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 44
ctggatectg ttgctagctt ttgaatgaaa 30

<210> 45
<211> 1050
<212> DNA
<213> Lycopersicon esculentum

<400> 45
aagcttatgg acagactaaa acactttttt ttttaataaa tattgtttgc aagtgtacac 60
cgaaagatct acgttattat aacataatat tacggtaaa gctagaagtc taattacgaa 120
tttcatgaga tttataact tttttttt ttatatttat attaaaaag tattaaatat 180
atacaaattt aaactcttaa aaccattgtt aaaaaattta gaatccaaaa tgtaatatt 240
atggtttgc ctctgctaaa cattactaat caaaaattatc tttttgttta gagcattact 300

24

actgtacaaa tctaccaagt ataaatataa aagctgttaa agaattccc cacacttatt	360
attcttaatc ttccacctac ccaatcacaa atatattaaa tgagcctcta aatttgcct	420
attgcgggta atatgatcta cctatcaatt atttgaatc tagtcaaaaa gatgccaaaa	480
aaatataata ctccatctag attgaaaatt tttgtcaata gaaaagaaga gaaacatgat	540
aactttataa aatattttac ctctggata gtttgatag acgtataat aataatata	600
taatattaat aaatgatgag attagttatc tttagaatgc attctatctt atgtctggtt	660
tgtatgtatta atgacaattt tgtttctaca accatgcatt attactgatc aatgtattgt	720
taaatgctaa tacgttgatt tgttatgtat tagttacata tacctatatg ttttgcata	780
agaaaaaatga tgtataacta attaataagt agtattatca tgtagtaagt tattttctg	840
gtcagtagag agcttctaag aacaaaaact aaataattgt attgtatggc tgctattcaa	900
aattccccac ctaacgcgtc ctggataat tgatatgact tgaagccgcc tctaaaatta	960
aataatattt ggtgcttata atgaaaaatac tattatataa agcaaggat agcccaatga	1020
attttcattc aaaagcttagc aacaggatcc	1050

<210> 46
<211> 25
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 46
aaaatttggg atctagaagg tgagg

25

<210> 47
<211> 22
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 47
acatgaaact ttgaatgctt tg

22

<210> 48
<211> 754
<212> DNA
<213> Lycopersicon pennellii

<400> 48
tctagaaggt gaggaacttt ttttaacaat atataagtaa gcattggta taatttcaca 60
acaacattac ggtaaaacct ctataaatta atacccgata aattaataat ccctctaaaa 120
taatattttt ctaggatttt cgattaggc aatgaaaaaaaa atcaccattt tcaataaaaat 180
aatgagataa tatattttca gaagaccctt atataaatac atgggccta ttaatatcat 240
aaattgatta ttattcaaaa gcataaataat atctaagata atttagtaaa aaaatgattc 300
tattctgttt tttttttgtt aaaatttaaa tgtagttgaa gttcatttct aacatttcat 360
attgcttcca agagctccaa ttttgtcttt tcgaacttca ccatagaaga gttccagatg 420
cgataagtgt ttccttacgc gtaactgggtt ccaaagttat agtacatcatat tcaacttcat 480
catcgacatt gcttttccg atggtatcca taaattcttc taagcttatt tgaaatggag 540
taatatttta tttggccccca acacattata taaggcaatg tatagcccta tgaatcttcg 600
tccaaagcta gcaataatgs caagtttgtg taatagtagt agtacatctc tcaaaactcc 660
tttacttct tccaccactt gtttatcttc cactcctaag ccctctcaac tttcctaca 720
tgaaaaacgt aacaaagcat tcaaagtttc atgt 754

<210> 49
<211> 30
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 49
atggaaaagc ttatggacag actaaaaacac 30

<210> 50
<211> 30
<212> DNA

<213> Artificial sequence

<220>

<223> single strand DNA oligonucleotide

<400> 50

ttccccgggac atgaaaacttt gaatgccttg 30

<210> 51

<211> 1179

<212> DNA

<213> Lycopersicon esculentum

<400> 51

aagcttatgg acagactaaa acactttttt ttttaataaa tattgtttgc aagtgtacac 60

cgaaaagatct acgttattat aacataatata tacggtaaa gctagaagtc taattacgaa 120

tttcatgaga tttataact tttatTTTA ttatattttt atttaaaaag tattaaatata 180

atacaaattt aaactcttaa aaccattgtt acaaaattta gaatccaaaa tgttaatatt 240

atggTTTcgc ctctgctaaa cattactaat caaaatttac tttttgttta gagcattact 300

actgtacaaa tctaccaagt ataaatataa aagctgttaa agaatttccc cacacttatt 360

attcttaatc ttccac^ctac ccaatcacaa atatattaaa tgagcctcta aatttgcct 420

attgcgggta atatgatcta cctatcaatt atttgtatc tagtcaaaaa gatccaaaa 480

aaatataata ctccatctag attgaaaatt tttgtcaata gaaaagaaga gaaacatgat 540

aactttataa aatattttac ctctggtata gttttgatata agcgtataat aataatata 600

taatattaat aaatgatgag attagttatc tttagaatgc attctatctt atgtctggtt 660

tgatgtatta atgacaattt tgTTTctaca accatgcatt attactgttc aatgtattgt 720

taaatgctaa tacgttgatt tgTTTatgtat tagttacata tacctatatg ttttgtata 780

agaaaaaatga tgtataacta attaataagt agtattatca tgagtaaagt tattttctg 840

gtcagtagag agcttctaag aacaaaaact aaataattgt attgtatggc tgctattcaa 900

aattccccac ctaacgcgtc ctggaataat tgatatgact tgaagccgca tctaaaatta 960

aataatattt ggtgcttata atgttttaca tattatataa agcaaggtat agccaaatga 1020

atTTTcatTC aaaagcttagc aacaatggca agtttgcgtta gtaatagtag tactacttct 1080

ctcaaaaactc ctttcacttc tttaggttcc actccaaagc cttgtcaact tttcctacat 1140

ggaaaacgta acaaaggcatt caaagttca tgtccccggg 1179

<210> 52

<211> 30

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 52

atggaaaagc ttatggacag actaaaaacac 30

<210> 53

<211> 26

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 53

aaccggggag ccgatgcagc taatgg 26

<210> 54

<211> 1299

<212> DNA

<213> Lycopersicon esculentum

<400> 54

aagcttatgg acagactaaa acactttttt ttttaataaa tattgtttgc aagtgtacac 60

cgaaaagatct acgttattat aacataatat tacgggtaaa gctagaagtc taattacgaa 120

tttcatgaga tttaataact tttatTTTA ttatTTTAT atTTAAAAG tTTAAATAT 180

atacaaattt aaACTCTTAA aaccATTGTT ACAAAATTAA gaATCCAAA TGTTAATATT 240

atggTTTcgc ctctgctaaa cattactaat cAAAATTATC TTTTGTTA gagcattact 300

actgtacaaa tctaccaagt ataaatataa aagctgttaa agaattccc cacacttatt 360

attcttaatc ttccacctac ccaatcacaa atatattaaa tgagcctcta aatttgcct 420

28	
attgcgggta atatgatcta cctatcaatt atttgaatc tagtcaaaaa gatgccaaa	480
aatataata ctccatctag attgaaaatt tttgtcaata gaaaagaaga gaaacatgat	540
aactttataa aatattttac ctctggata gttttgatat agcgataat aataatata	600
taatattaat aaatgatgag attagttatc tttagaatgc attctatctt atgtctggtt	660
tgatgtatta atgacaattt tgtttctaca accatgcatt attactgatc aatgtattgt	720
taaatgotaa tacgttgatt tgttatgtat tagttacata tacctatatg ttttgtataa	780
agaaaaatga tgtataacta attaataagt agtattatca tgagtaaagt tattttctg	840
gtcagtagag agcttctaag aacaaaaact aaataattgt attgtatggc tgctattcaa	900
aattccccac ctaacgcgtc ctggaataat tgatatgact tgaagccgac tctaaaatta	960
aataatattt ggtgcttata atgtttaca tattatataa agcaaggtat agccaaatga	1020
attttcattc aaaagctagc aacaatggca agtttgtgt aataatagtag tactacttct	1080
ctcaaaactc ctttcaattc tttaggttcc actccaaagc cttgtcaact tttcctacat	1140
ggaaaacgta acaaaggcatt caaagttca tgcaaggtt ccaataactaa cggttaaccaa	1200
gatgaaacga attctgtaga tcgaaggaat gtttttcttg gcttagggagg tctttatggt	1260
gttgctaatg ctataccatt agctgcattcg gtcgggg	1299

<210> 55
<211> 25
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 55
gggatgagct acaacttgct tggat

25

<210> 56
<211> 23
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 56

ctaggagctc ttcagttcg gag

23

<210> 57

<211> 511

<212> DNA

<213> Homo sapiens

<400> 57

gggatgagct acaaacttgc tggattccta caaagaagca gcaatttca gtgtcagaag 60
 ctccctgtggc aattgaatgg gaggcttgaa tattgcctca aggacaggat gaactttgac 120
 atccccctgagg agattaagca gctgcagcag ttccagaagg aggacgcgc attgaccatc 180
 tatgagatgc tccagaacat ctttgcattt ttcagacaag attcatctag cactggctgg 240
 aatgagacta ttgttgagaa cctccctggct aatgtctatc atcagataaa ccatctgaag 300
 acagtccctgg aagaaaact ggagaaaagaa gattttacca ggggaaaact catgagcagt 360
 ctgcacctga aaagatatta tgggaggatt ctgcattacc tgaaggccaa ggagtacagt 420
 cactgtgcct ggaccatagt cagagtggaa atcctaaggaa acttttactt cattaacaga 480
 cttacagggtt acctccgaaa ctgaagagct c 511

<210> 58

<211> 592

<212> DNA

<213> Artificial sequence

<220>

<223> Synthetic human growth hormone gene

<400> 58

cccgggatgt tcccaactat tccattgtct aggctttcg ataatgtat gttgagggt 60
 catagggttgc atcagttggc tttcgataact taccaggagt tcgaggaggc ttacattcca 120
 aaggaggcaga agtactcatt ctttcagaat ccacagactt ctttgcgtt ctctgagtct 180
 attccaaactc catcaaatacg ggaggagact cagcagaagt caaatcttga gttgttgg 240
 atttctttgt tgcttattca gtcttgggtt gagecagttc agttcttgag gagtgttttc 300
 gcaaattctt tggtttacgg agcttcagat tcaaatagttt acgatttggta gaaggatttg 360

30

gaggagggaa ttcagactct tatgggaagg ttggaggatg gatctccaag gactggacag	420
attttcaagc agacttactc taagttcgat acaaactctc ataacgatga tgctttgttg	480
aagaattacg gattgttcta ctgcttcagg aaggatatgg ataaggttga gactttcttg	540
aggattgttc agtgcaggag tggtgaggga tcttgeggat tctgatgac tc	592

<210> 59

<211> 252

<212> DNA

<213> Artificial sequence

<220>

<223> PPOD lumen signal peptide coding sequence

<400> 59

atggcaagtt tgtgttagtaa tagtagtact acttctctca aaactccctt cacttcttta	60
ggttccactc caaagccttg tcaacttttc ctacatggaa aacgtaacaa agcattcaaa	120
gtttcatgca aggttaccaa tactaacggt aaccaagatg aaacgaattc tgtagatcg	180
aggaatgttc ttcttggctt aggaggtott tatggtgttg ctaatgctat accattagct	240
gcatcggttc cc	252

<210> 60

<211> 84

<212> PRT

<213> Artificial sequence

<220>

<223> PPOD lumen signal peptide sequence

<400> 60

Met Ala Ser Leu Cys Ser Asn Ser Ser Thr Thr Ser Leu Lys Thr Pro			
1	5	10	15

Phe Thr Ser Leu Gly Ser Thr Pro Lys Pro Cys Gln Leu Phe Leu His		
20	25	30

Gly Lys Arg Asn Lys Ala Phe Lys Val Ser Cys Lys Val Thr Asn Thr		
35	40	45

Asn Gly Asn Gln Asp Glu Thr Asn Ser Val Asp Arg Arg Asn Val Leu
50 55 60

Leu Gly Leu Gly Gly Leu Tyr Gly Val Ala Asn Ala Ile Pro Leu Ala
65 70 75 80

Ala Ser Ala Pro

<210> 61
<211> 141
<212> DNA
<213> Artificial sequence

<220>
<223> PPOA stromal signal peptide coding sequence

<400> 61
atggcaagtt tgtgttaatag tagtagtaca tctctcaaaa ctcctttac ttcttccacc 60
acttgttat cttccactcc taagccctct caactttcc tacatggaaa acgtaacaaa 120
gcattcaaag tttcatgtgg g 141

<210> 62
<211> 47
<212> PRT
<213> Artificial sequence

<220>
<223> PPOA stromal signal peptide sequence

<400> 62

Met Ala Ser Leu Cys Asn Ser Ser Ser Thr Ser Leu Lys Thr Pro Phe
1 5 10 15

Thr Ser Ser Thr Thr Cys Leu Ser Ser Thr Pro Lys Pro Ser Gln Leu
20 25 30

Phe Leu His Gly Lys Arg Asn Lys Ala Phe Lys Val Ser Cys Gly
35 40 45

<210> 63
<211> 261
<212> DNA
<213> Artificial sequence

<220>
<223> PPOA lumen signal peptide coding sequence

<400> 63
atggcaagtt tggtaatag tagtagtaca tctctaaaaa ctcctttac ttcttccacc 60

acttgtttat cttccactcc taagccctct caactttcc tacatggaaa acgtAACAAA 120

gcattcaaag tttcatgtgg gaaggttacc aatactaacg gtaaccaaga tgaaacgaat 180

tctgttgatc gaagaaatgt tcttcttggc ttaggtggc tttatggtgt tgctaatgtc 240

ataccattag ctgcattccgc t 261

<210> 64
<211> 87
<212> PRT
<213> Artificial sequence

<220>
<223> PPOA lumen signal peptide sequence

<400> 64

Met Ala Ser Leu Cys Asn Ser Ser Ser Thr Ser Leu Lys Thr Pro Phe
1 5 10 15

Thr Ser Ser Thr Thr Cys Leu Ser Ser Thr Pro Lys Pro Ser Gln Leu
20 25 30

Phe Leu His Gly Lys Arg Asn Lys Ala Phe Lys Val Ser Cys Gly Lys
35 40 45

Val Thr Asn Thr Asn Gly Asn Gln Asp Glu Thr Asn Ser Val Asp Arg
50 55 60

Arg Asn Val Leu Leu Gly Leu Gly Leu Tyr Gly Val Ala Asn Ala
65 70 75 80

Ile Pro Leu Ala Ala Ser Ala

85

<210> 65

<211> 111

<212> DNA

<213> Artificial sequence

<220>

<223> Plastid signal peptide, predicted to direct protein to the stroma

<400> 65

gaccgcctcca ctcccaaaaa caacacaccaa tattcaagga tgatagttgg ctatagaagc 60

acaatcatta ccctttctca tccttaagcta ggcaatgggg aaacaatttc a 111

<210> 66

<211> 138

<212> DNA

<213> Artificial sequence

<220>

<223> Plastid signal peptide, predicted to direct protein to the stroma

<400> 66

atgagttttt tggtttttca atgttggaaa ttatcatctc catctctgtat tttacaacaa 60.

aatacatcaa tatccatggg tgcattcaaa ggtattcata aacttcaaat cccaaattca 120

cctctgacag tgtctgt 138

<210> 67

<211> 1834

<212> DNA

<213> Nicotiana tabacum

<400> 67

tcttagactac catcgctagt aatcggtggca ataactaccc taactatagc atttattgtc 60

accaaataaa atttggcagc taatcataat tttttgtcat gaatcaatag ttattgtgc 120

aatagttatc tcttagccac aataaaattat ttaaaaataaa atattatagc taaaataaata 180

tttttgcttt aagttctaaa agcttgtggc aatagttaaa tgatatagtc acagatttat 240

34

tggtataatt	gaattatgtt	gctaatttct	tagtttttg	ccacgagtta	aaaattacca	300
atagctatag	taactttta	atcacaataa	aatatttcaa	agaaaatatt	gtagctaaat	360
gaatattttt	tccttcaagt	tataaaagt	tgtggcaata	taggttaat	tagccacatg	420
tttcttgctt	taatagaatt	ttttagctaa	tcattaactt	ttaccacgag	ttgaacttaa	480
tataacaaca	ataacctttt	aaccataata	aagcgattta	aatcaaataat	tactaaataa	540
ataactttgc	tttcaagttt	ctataaaatc	atggcaatag	tcattacgat	aaaatgatat	600
aaccacgaat	atattgcaac	gataaattct	gtaactaatac	attagttttt	gcgacgaggt	660
aaattttccg	tcacagtgc	aatcttctag	gcacattaaa	aatttcaaac	aaaattttgt	720
agtcaaataa	atatttatct	tcttattttt	agaaaataaa	aatagttaga	taatagttac	780
tactatttgt	catgaaaata	tcaatagata	caaattttaa	gtgactataa	atttacgagt	840
ttactatact	ttagtcgtac	agtttgcatt	aatagtattt	taaccacaat	tagttatag	900
tacaaaataa	cataagtgaa	taactttttt	tcaatgagaa	aataagagtt	gctcaaacaa	960
tatcaaggta	caaaaattta	attttaactg	aaaaagttat	attttccaa	aataacataa	1020
actatagtaa	ttatatatag	tttjaagtat	taataaaatt	taaatatgca	aaagttaatt	1080
ttaataaacc	atttgtatgc	ctaacttgta	gcctctaaac	tattttattt	gctttattta	1140
tcaaactcat	attttatttt	attgcacctt	gttagtttg	gacgttaatt	atatatattt	1200
ggtgtaaaat	ttaaaatata	ttaacatttg	tggagaattt	atgtatgcct	ggttcttaac	1260
tatttttttt	tatataactg	gttagagtaa	tttcttatat	ttcagttat	attttaaat	1320
aagtcctcat	aaattgaaga	ctttaaaagt	ttttgtgtca	ttcctttttt	tatthaagaa	1380
attgaagaat	tccgctaaat	ttcatatttc	cgctgttatt	taactgttta	tttccccttgt	1440
taatataatt	ggtaagaagt	tttaaaataa	aggagttaat	gatttcttag	gttcatggct	1500
tgcctagctt	ctacgagtaa	gcccacatcac	gactccccag	gataaggaaa	tccgggtcgt	1560
agcattcact	cacaaaaatt	actaaaaaca	aagtttaccc	ttctccaaa	agtaaatttc	1620
atatttggct	ccacataatg	tgttcaatga	gtcaagtgaa	gtactttca	tgacaazzaaa	1680
aagttgctga	aaaatgcata	tctcatattt	ttttttttaga	gaaatcccat	ttcttgctta	1740

35

aacgaaagcc tataaaagag catatattgc aacaacagt tgcagaaaact atcaagtcaa	1800
ataatcccc ctttaattcc ctcccaaacc cggg	1834
<210> 68	
<211> 1969	
<212> DNA	
<213> Nicotiana tabacum	
<400> 68	
tctagataag ttgataaagc taatttctca ttttagctac catcgctagt aatcggtggca	60
ataaactaccc taactatagc atttatttgct accaaataaa atttggcagc taatcataat	120
tttttgtcat gaatcaatag ttattgttagc aatagttatc tcttagccac aataaatttat	180
ttaaaataaa atattatagc taaataaataa tttttgtttt aagttctaaa agcttgtggc	240
aatagttaaa tgatatagtc acagatttat tggtataatt gaattatgtt gctaatttct	300
tagtttttg ccacgagttttaaaaattacca atagctatag taactttta atcacaataa	360
aatattttgaa agaaaaatattt gtagctaaat gaatattttt tccttcaagt tattaaaagt	420
tgtggcaata taggttaaat tagccacatg tttttgtttttaatagaatt ttgttagctaa	480
tcattaactt ttaccacgag ttgaacttaa tataacaaca ataaccctttt aaccataata	540
aagcgattta aatcaaataat tactaaataa ataactttgc tttcaagttt ctataaaatc	600
atggcaatag tcattacgat aaaatgatat aaccacgaat atattgcaac gataaattct	660
gtactaatac attagttttt ggcacgaggtaaattttccgtc acagtagc aatcttcttag	720
gcacattaaa aatttggaaac aaaattttgtt agtcaaataa atatttatct tcttattttta	780
agaaaataaa aatagttttaa taatagttac tactattttgtt catgaaaataa tcaatagata	840
caaattttaaa gtgactataa atttacgagt ttactataact ttagtcgtac agtttgcaat	900
aatagttttt taaccacaat tagttatag tacaaaataa cataagtgaa taactttttt	960
tcaatgagaa aataagagtt gctcaaacaa tatcaagttt caaaaattta attttaactg	1020
taaaagttat atttttccaa aataacataa actatagtaa ttatataatag tttgaagtat	1080
taataaaaattt taaatatacgaa aaagttttaattt ttaataaacc atttgcgtac ctaacttgta	1140
gcctctaaac tatttttattt gcttttattta tcaaactcat atttttattt attgcacattt	1200

gttagtttg gacgtaatt atatatattt ggtgtaaaat ttaaaatata ttaacattt	1260
tggagaattt atgtatgcct ggttcttaac tattttttt tatataactg gttagagtaa	1320
tttccttat ttcagtattt atttttaat aagtcctcat aaattgaaga cttaaaagt	1380
ttttgtgtca ttccctttt tatttaagaa attgaagaat tccgctaaat ttcatattc	1440
cgtgttatt taactgttta ttcccctgt taatataatt ggtaagaagt tttaaaataa	1500
aggagttaat gattttctag gttcatggct tgccctagctt ctacgagtaa gcgccatcac	1560
gactcccgag gataaggaaa tccgggttgt agcattcaact cacaaaaatt actaaaaaca	1620
aagtttaccc ttctccaaa agtaaatttc atatttggct ccacataatg tttcaatga	1680
gtcaagtgaa gtactttca tgacaaaaaa aagttgctga aaaatgcata tctcatattt	1740
tttttttaga gaaatcccat ttcttgctta aacgaaagcc tataaaagag catatattgc	1800
aacaacagtt tgcagaaact atcaagtcaa ataatcccc ctttaattcc ctccaaaaat	1860
gcagttcttc aacttctttt ccctttctt tttgtgtca tttcttttt tatttaagaa	1920
atgaaagaat tccaatagcc aaacccaaag attgcctcca gttcccccggg	1969

<210> 69
<211> 806
<212> DNA
<213> Lycopersicon esculentum

atgaaagtaa ctttggta tagtacttca ctctctattt tgtttggct tctacttgtt	60
aaacttgttt catcaaaacg aagaaaacag aatctaccac caagcccaact acttaaactt	120
ccaatattag gccatctcta tctccttaaa ccacmtctat atcgactct tgctaatctc	180
tcaactaaat atggccctgt tttctcttt caatttagta cccgtttgt tgtagcaatt	240
tcctcaccat ctgctgccga agaatgttcc acaaaaaaaaatg atatcgttt tgctaatcgc	300
cctcggacaa tgacggaaaa attcataggc tataactcta ctacagtcat tggttctcct	360
tatggtgatc actggcgctt ctctggccgc ctctgcccac ttgaaatatt ctccactaat	420
cgtctcaaca attttcagtc cattagacaa gatgaaatca aacttttagt tcgaagagtg	480

37

tttcacaat ctggagacaa ttttgtgact cctgttgagc ttaagtccaa gctttttcag	540
atgtcgata atattatcat gagaatggta gctggaaaaa gatattacgg tgaagagata	600
gataacgagg aggcaaatca ttttgggtg ctttagaag argttatttc ktttgggggt	660
gtatcaaatg ccghggattt catgcctgca atatttctgk tgttttcag gagtacggag	720
aaaaaaaaatag caaagcttgg taataagatg gacaagstct tgcaaggttt ggktgatgaa	780
catcgccgca ataaaagcag gaatac	806
<210> 70	
<211> 912	
<212> DNA	
<213> Lycopersicon esculentum	
<400> 70	
atgagttctt tggttttca atgttggaaa ttatcatctc catctctgat tttacaacaa	60
aatacatcaa tatccatggg tgcattcaaa ggtattcata aacttcaaat cccaaattca	120
cctctgacag tgtctgctcg tggactcaac aagatttcat gctcactcag cttacaaacc	180
aaaaaaacttt gttatgagga taatgataat gatcttgatg aagaacttat gcctaaacac	240
attgcttga taatggatgg taataggaga tggcaaaagg ataagggttt agacgtatecc	300
gaagggtcaca aacatctttt tccaaaatta aaagagattt gtgacatttc ttctaaattg	360
ggaatacaag ttatcactgc ttttgcattc tctactgaaa attggaaacg agccaagggg	420
gaggttgatt tcttgatgca aatgttgcggaa gaactctatg atgagtttc gaggtctgga	480
gtaagagtgt ctattattgg ttgtaaaaco gacccccaa tgacattaca aaaatgcata	540
gcattaacag aagagactac aaagggaaac aaaggacttc accttgcatttgcactaaac	600
tatggtggat attatgacat attgcaagca acaaaaagca ttgttaataa agcaatgaat	660
ggtttattag atgtagaaga tatcaacaag aatttatttg atcaagaact tggaaagcaag	720
tgtccaaatc ctgatattact tataaggaca ggaggtgatc aaagagtttag taactttttg	780
ttgtggcaat tggcttayac tgaattttac ttcaccaama cattgtttcc tgattttgga	840
gaggaagatc taaaagaggc aataatraac tttcaacaaa ggcatacgtacg ttttgggtgga	900
cacacatattt ga	912

<210> 71
 <211> 1269
 <212> DNA
 <213> Lycopersicon esculentum

<400> 71	
atgggtaata ttcatttgat ttcccactt ttatttatat cttgtttcat tttcccatcc	60
acaacaaatg gctactccaa cgcaatcata aaagetttgt gcacccaaac acctcatcca	120
caaccttgtg aatacttctt atcacaaaat cccaaaatta catctcctat cataaaaaaa	180
tcagatttgc taaaagtgtc actagactta gtgttagacc gtgcgttacg tgcccaactg	240
aacacatatt cactaggc aaaaatgtcg aacgagcgcg aaaaaaacgc atgggctgat	300
tgcattgaac tctatgaaaa ctcacatcaac aaaatcaaaa gcacagttga tccaaacaca	360
aatgctcag ctactgatgc tcaaacatgg ttaagtacat ccttaacaaa tcttgaacaca	420
tgtaaagcag gtttgcaga attaggcggtt acggattatg ttatgccact aatatcaaata	480
aataatgtgt catcttaat aagtaacgct ttagcttta atcatggta ttatactgaa	540
cctactaaaa gtagtactac tactcaagtt gatggatttc caacttgggt atctcctgg	600
gatagaaaaat tggcaatc gtcgcgtcg tggcgatcaa cggcttctca ggcaatgt	660
gtggggcata ctgatggttc aggggatttt aagacagtga aagaagctgt agatgctgct	720
gccaagaata aaggaagtgg gaggtttgtg atatatgtga aagctggac ttataatgaa	780
aatgtggaga ttggagaaaa ggtgaaaaat gttatgtga ttggagatgg cattggaaag	840
acaattatta ctggaaagcaa aagtgttgg agtggatcca ccacctttag atcagccaca ,	900
gttggtgctt ctggtgacgg atttattgtc caaggcataa caattagaaa cactgctgga	960
ccccaaaagc accaagcagt agccctacga tctggctctg atcttcagt attttatcaa	1020
tgtagcttcg aagggtatca agacactttg tacgttcatt ccaataggca attttacaaa	1080
gagtggtgata ttatgttac ggtcgatttt atatggtm acgcagcagt tgtattacaa	1140
aattgtata tttcgcttag agaccctccg aataaaaatca acactgtgac agcccaaggg	1200
cgaaccggacc cgaatcaaaa cactggatttccatata attgtagaat cactggatc	1260

40

gctagagttt	ctctacagag	agatgtatgtt	gaacttagtt	cagtgaatga	tccatttatt	120
tccactgatt	acatgacata	tatgttttaag	tatgatttcag	tacatggaca	atgaaagcat	180
catgagctaa	aggtaaggaa	tgagaagaca	cttcttttgc	gagagaaggc	tgttacagtt	240
tttggaatca	ggaaccctga	agatatccca	tggggtgaag	ctgggtgtga	cttcgttgtt	300
gaatcaaccg	gtgtcttcac	tgacaaggac	aaggctgtcg	ctcaacttcaa	gggtgggtgc	360
aagaaggttt	tgtatctgtc	tcttagcaaa	gatgtccca	tgtttgttgt	gggtgtcaac	420
gagaatgaat	acaaggccaga	gctggacatt	gtctccaatg	ctagttgcac	aacgaactgc	480
cttgcacctt	tggctaagggt	tatcaatgtat	aggtttggca	ttgttgagggt	tctcatgacc	540
actgtccacg	ccatgactgc	cacccagaaa	actgttgatg	gtccatccat	gaaggactgg	600
agaggtggaa	gagctgttcc	attcaacatc	atccctagca	gcactgggtgc	agccaaggct	660
gttggaaaag	tgtcccaca	acttaacggc	aaattgactg	aatggcctt	cagagtacca	720
actgctgatg	tctccgttgt	cgatcttact	gtaagactcg	agaaagaagc	ctcctatgaa	780
gacattaagg	ctgcaatcaa	ggaggaatca	gagggttaat	tgaagggttat	cttggataac	840
actgaagatg	atgtggtttc	cacagacttt	gttggtgaca	gcaggtcaag	catttttgat	900
gccaggctg	gaattgtttt	gagcaagaat	tttggaaag	ttgtgtcatg	gtatgacaac	960
gaatggggtt	acagttcccc	tgtgattgtat	ttgtatctgc	atatggctaa	ggcttga	1017

<210> 74
<211> 657
<212> DNA
<213> Lycopersicon esculentum

<400> 74

atgggtgtcac	tgaaacttca	gaagcggctc	gcgcgcagtg	ttctaaagtgc	tgggggggaa	60
aaagtatggc	ttgaccctaa	cgaaggcaat	gaaatctcca	tggcttaactc	aaggcaaaac	120
atcagaaaatgt	tggtaagggaa	tggtttcatc	atcaggaaac	caaccaaaat	tcactcacga	180
tctcggtcac	gcaggatgaa	ggaagccaaa	agaaaggcc	gtcactctgg	atatggtaag	240
cgtaagggttac	ccagggaggc	tagttgccc	acaaagggtgc	tgtggatgag	gagactcaga	300
gtcctcaggc	gtttgtttcg	taagtacagg	gagtccaaaga	agattgacaa	gcacatgtac	360

41

catgatatgt acatgaaggt gaagggtaat gtcttcaaga acaagcgtgt tctcatggag	420
aacattcaca aaaccaaggc tgagaaggct agagagaaga cttgtctga ccaattttag	480
gccaggaggg caaagaacaa ggcaagcagg gaaagaaaagt tcgttaggag ggaggaacgt	540
ttggcccgagg gaccaggaga gaagccagta caacctgcag cgccagcccc ggccaccagca	600
gcaacagcac ccccagccaa gactgctcag ggaggatcta agaagtcaaa gaagtga	657

<210> 75
<211> 132
<212> DNA
<213> Artificial sequence

<220>
<223> PPOD stromal signal peptide coding sequence

<400> 75	
atggcaagtt tgtgtagtaa tagtagtact acttctctca aaactccttt cacttcttta	60
ggttccactc caaaggccttg tcaacttttc ctacatggaa aacgtAACaa agcattcaaa	120
gtttcatgtc cc	132

<210> 76
<211> 44
<212> PRT
<213> Artificial sequence

<220>
<223> PPOD stromal signal peptide coding sequence
<400> 76

Met Ala Ser Leu Cys Ser Asn Ser Ser Thr Thr Ser Leu Lys Thr Pro			
1	5	10	15

Phe Thr Ser Leu Gly Ser Thr Pro Lys Pro Cys Gln Leu Phe Leu His		
20	25	30

Gly Lys Arg Asn Lys Ala Phe Lys Val Ser Cys Pro	
35	40

42

<210> 77
<211> 20
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 77
ccacatgccca ttctccgtct

20

<210> 78
<211> 21
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 78
gcttttcttt cacgtccctg a

21

<210> 79
<211> 21
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 79
ttgttgtggg tgtcaacgag a

21

<210> 80
<211> 19
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 80
atggcgtgga cagtggta

19

<210> 81
<211> 25

43

<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 81
caactctggat atggtaagecg taagg

25

<210> 82
<211> 24
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 82
ttctttggact ccctgtactt acga

24

<210> 83
<211> 25
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 83
tctcttcaat taggtacccg tcttg

25

<210> 84
<211> 21
<212> DNA
<213> Artificial sequence

<220>
<223> Single strand DNA oligonucleotide

<400> 84

tgaattttgc cgtcattgtc c

21

<210> 85
<211> 23
<212> DNA
<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 85

gggttttagac gtatccgaag gtc

23

<210> 86

<211> 25

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 86

gctcgttcc aattttcagt agaga

25

<210> 87

<211> 21

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 87

ttacgtgccc aactgaacac a

21

<210> 88

<211> 19

<212> DNA

<213> Artificial sequence

<220>

<223> Single strand DNA oligonucleotide

<400> 88

caatgcaatc agcccatgc

19

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
23 December 2004 (23.12.2004)

PCT

(10) International Publication Number
WO 2004/111183 A3

(51) International Patent Classification:
CI2N 15/00 (2006.01) *C07H 21/00* (2006.01)
CI2N 15/82 (2006.01)

(81) Designated States (*unless otherwise indicated, for every kind of national protection available*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW

(21) International Application Number:
PCT/IL2004/000549

(22) International Filing Date: 20 June 2004 (20.06.2004)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
60/479,467 19 June 2003 (19.06.2003) US

(71) Applicant (*for all designated States except US*): EVO-GENE LTD. [IL/IL]; 13 Gad Finstein Street, 76 121 Rehovot (IL).

(72) Inventors; and

(75) Inventors/Applicants (*for US only*): RONEN, GU [IL/IL]; 20 Moshav Ometz, 33 870 Emek Hefer (IL). RABINOVICH, Larisa [IL/IL]; 13/13 Hanagid Street, 75 482 Rishon LeZion (IL). MEISSNER, Rafael [IL/IL]; 11 Halevona Street, 76 350 Rehovot (IL). KARCHI, Hagai [IL/IL]; Moshav Sitiya, 76 834 Moshav Sitiya (IL).

(74) Agent: G.E. EHRLICH (1995) LTD.; 11 Menachem Begin Street, 52 521 Ramat Gan (IL).

(84) Designated States (*unless otherwise indicated, for every kind of regional protection available*): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

- with international search report
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments

(88) Date of publication of the international search report:
16 March 2006

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

WO 2004/111183 A3

(54) Title: PLANT TRICHOME-SPECIFIC PROMOTER AND LEUCOPLAST SIGNAL SEQUENCE

(57) Abstract: Novel plant-derived regulatory sequences and constructs and methods of using such sequences for directing expression of exogenous polynucleotide sequences in trichomes are provided. In addition, a polynucleotide capable of targeting a protein to leucoplasts is provided. Methods for the heterologous expression of proteins are disclosed. Methods including the reduction of endogenous polyphenol oxidase (PPO) enzyme are disclosed.

INTERNATIONAL SEARCH REPORT

International application No

PCT/IL04/00549

A. CLASSIFICATION OF SUBJECT MATTER

IPC(8) C12N 15/00, 15/82; C07H 21/00
 US CL 800/278, 287, 536/23 1, 24.1; 435/468, 419

According to International Patent Classification (TPCI or to both national classification and IPC)

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
 U.S. 800/278, 287; 536/23.1, 24.1; 435/468, 419

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
 Please See Continuation Sheet

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
A	LARKIN, J.C et al. Arabidopsis GLABROUSI Gene Requires Downstream Sequences for Function. Plant Cell 1993, Vol 5 No. 12, pages 1739-1748, entire document	1-17, 19-21, 23-27, and 29-30
A	HEDTKE, B et al. Green fluorescent protein as a marker to investigate targeting of organellar RNA polymerases of higher plants in vivo. Plant Journal 1999, Vol. 17, No. 5, pages 557-561, entire document	1-17, 19-21, 23-27, and 29-30
A	MENZEL, G. et al. Expression of bacterial poly(3-hydroxybutyrate) synthesis genes in hairy roots of sugar beet (Beta vulgaris L.). Appl. Microbiol. Biotechnol. 2003 Vol. 60, pages 571-576, entire document.	1-17, 19-21, 23-27, and 29-30

 Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"A" document defining the general state of the art which is not considered to be of particular relevance	"X"	document of particular relevance, the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"E" earlier application or patent published on or after the international filing date	"Y"	document of particular relevance, the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"&"	document member of the same patent family
"O" document referring to an oral disclosure, use, exhibition or other means		
"P" document published prior to the international filing date but later than the priority date claimed		

Date of the actual completion of the international search

06 January 2006 (06.01.2006)

Date of mailing of the international search report

06 JAN 2006

Name and mailing address of the ISA/US

Mail Stop PCT, A/H: ISA/US
 Commissioner for Patents
 P.O. Box 1450
 Alexandria, Virginia 223 13-1450
 Facsimile No. (571) 273-3201

Authorized officer

Cathy K. Worley

Telephone No. (571) 272-0500

INTERNATIONAL SEARCH REPORT

International application No.

PCT/IL04/00549

Box No. π Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely.

2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. I Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. DI Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:
Please See Continuation Sheet

1. I As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying additional fees, this Authority did not invite payment of any additional fees.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

4. X J No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: Please See Continuation Sheet

Remark on Protest

<input type="checkbox"/>	The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.
<input type="checkbox"/>	The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
<input type="checkbox"/>	No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/IL04/00549

Continuation of Item 4 of the first sheet:

The title is too long, PCT Rule 4.3. New title: "Plant trichome-specific promoter and leucoplast signal sequence".

BOX III. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

Groups 1-1200 contain claims directed to patentably distinct inventions. There are four separate instances of multiple versions being claimed: A) The polynucleotide capable of regulating expression, B) the nucleic acid sequence (presumably encoding a transit peptide), C) the small molecule capable of interfering with expression accumulation or stability of the polypeptide of interest, and D) the promoter utilized in gene silencing. Each invention in groups 1-1200 is drawn to a different combination of these four elements. The four elements are as follows:

- A) an isolated polynucleotide capable of regulating expression of at least one polynucleotide sequence operably linked thereto in trichomes comprising a nucleic acid sequence at least 80% identical to SEQ ID NO: 23, 26, or 29.
- B) a nucleic acid sequence selected from SEQ ID NOS: 59, 61, 63, 65, and 67.
- C) said at least one molecule being selected from: PPO enzyme, an enzyme, a metabolite, a polyphenol, a ketone, a terpenoid, a phenylpropanoid, and alkaloid.
- D) said method comprising gene silencing utilizing a nucleic acid comprising a nucleic acid sequence selected from SEQ ID NOS: 23, 26, 29, 35, 38, 39, 42, 48, 50, and 51.

In the absence of an election, the examiner will search and write an opinion for Group 1 which will comprise the first item in each of the four lists A-D.

If the applicant would like to choose one of the other groups selected from groups 2-1200, then the applicant must specify one sequence or molecule from each of the four lists below:

- For A) the applicant must elect one of the sequences selected from SEQ ID NOS: 23, 26, or 29.
- For B) the applicant must elect one of the sequences selected from SEQ ID NOS: 59, 61, 63, 65, and 67.
- For C) the applicant must elect one of the molecules selected from: PPO enzyme, an enzyme, a metabolite, a polyphenol, a ketone, a terpenoid, a phenylpropanoid, and alkaloid.
- For D) the applicant must elect one of the sequences selected from SEQ ID NOS: 23, 26, 29, 35, 38, 39, 42, 48, 50, and 51.

Group 1201, claims 21-22, drawn to a method of producing a molecule of interest in plant trichomes, the method comprising endogenously expressing a polypeptide capable of directly or indirectly increasing a level of the molecule of interest in the plant trichomes and down-regulating a level of at least one molecule endogenous to the plant trichomes, said at least one molecule being capable of interfering with accumulation or stability of the molecule of interest, thereby producing the molecule in the plant trichomes.

Group 1202-1241, claims 31-46, drawn to a plant genetically modified to express a molecule of interest in trichomes, wherein said plant is further modified or selected capable of accumulating less than 50% of average volume of undesired compounds in trichome cells of said plant species, and said plant comprising an expression construct, and said expression construct comprising a promoter selected from SEQ ID NOS: 23, 26, 29, 35, 38, 39, 42, and 45, and said expression construct additionally comprising a nucleotide sequence selected from SEQ ID NOS: 59, 61, 63, 65, and 67.

Groups 1202-1241 contain claims directed to patentably distinct inventions. There are two separate instances of multiple versions being claimed: E) a promoter and F) a nucleotide sequence. Each invention in groups 1202-1241 comprises a different combination of promoter and nucleotide sequence. The two lists are as follows:

INTERNATIONAL SEARCH REPORT

International application No.

PCT/IL04/00549

- E) a promoter selected from SEQ ID NOs: 23, 26, 29, 35, 38, 39, 42, and 45
- F) a nucleotide sequence selected from SEQ IDNOs: 59, 61, 63, 65, and 67

If the applicant chooses to elect one of the groups selected from groups 1202-1241, the applicant must further elect a sequence from E and F).

For E) the applicant must elect one of the promoters selected from SEQ ID NOs: 23, 26, 29, 35, 38, 39, 42, and 45.

For F) the applicant must elect one of the nucleotide sequences selected from SEQ IDNOs: 59, 61, 63, 65, and 67.

Group 1243, claims 47-51, drawn to a method of harvesting trichomes and/or exudates an/or content thereof, the method comprising: incubating a trichome-containing plant tissue in a liquid such that trichome exudates and content is released into said liquid, wherein incubating is effected while avoiding friction of said trichome-containing plant tissue with a solid surface; and collecting said liquid, to thereby harvest the trichome exudates and content.

Group 1243, claim 52, drawn to an apparatus for mechanical harvesting of trichome exudates and content, the apparatus comprising a mechanism designed and configured for mechanically agitating a trichome-containing plant tissue in a fluid and collecting said fluid to containing the trichome exudates or content.

According to PCT Rule 13.2, unity of invention exists only when there is a shared same or corresponding special technical feature among the claimed inventions, and said technical feature is a contribution over the prior art. The inventions listed as groups 1-1243 appear to share the common technical feature of being involved in the expressing or harvesting of a product in trichomes. The technical feature of expression in trichomes is shown by Larkin et al. (1993) Plant Cell Vol. 5 pp. 1739-1748 to lack novelty or inventive step. Larkin et al. teach that the promoter of the GL1 gene in conjunction with the 3' noncoding region direct expression of a GUS reporter gene in developing trichomes. Therefore the common technical feature of being involved in either the expressing or harvesting of a product in trichomes does not make a contribution over the prior art.

Continuation of Box III Item 4:

1-17, 19-21, 22-27, and 29-30 (claims 18 and 28 are drawn to non-elected subject matter of metabolites instead of PPO enzyme).

Continuation of B. FIELDS SEARCHED Item 3:

STIC Sequence search for SEQ ID NO:23 and SEQ ID NO:59.

STN search of the Agricola, Biosis, and Caplus databases; and WEST search were performed.

Search terms: trichomes, promoter, heterologous expression, transgenic, transgene, transformant, transform, leucoplast